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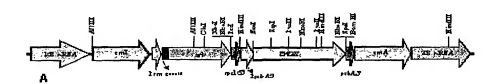
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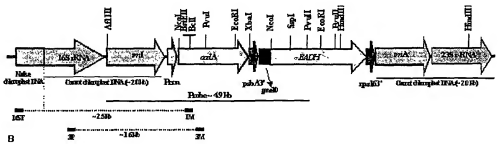
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(54) Title: PLASTID GENETIC ENGINEERING VIA SOMATIC EMBRYOGENESIS





(57) Abstract: A plastid transformation vector suitable for transforming a non-green plant cell, in which the plastid vector comprises, as operably linked components, a first flanking sequence, a DNA sequence coding for a foreign gene, and a second flanking sequence.

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For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

## PLASTID GENETIC ENGINEERING VIA SOMATIC EMBRYOGENESIS

# STATEMENT REGARDING FEDERALLY SPONSORED RESEARCH

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## FIELD OF THE INVENTION

The field of this invention relates to genetically engineering a plant plastid. More specifically, this invention relates throught the transformation of non-green plant cells through plastid transformation, and the subsequent regeneration the non-green plant cells through somatic embryogenesis.

## **BACKGROUND**

Plastids are ideal for genetic engineering because it offers a number of attractive advantages, including high-level transgene expression (Daniell et al., 2002), multi-gene engineering in a single transformation event (DeCosa et al., 2001; Ruiz et al., 2003; Daniell & Dhingra, 2002), transgene containment via maternal inheritance (Daniell 2002), lack of gene silencing (Lee et al., 2003; DeCosa et al., 2001), position effect due to site specific transgene integration (Daniell et al., 2002) and pleiotropic effects (Daniell et al., 2001; Lee et al., 2003). Chloroplast genetic engineering is most suitable for hyper-expression of vaccine antigens and production of valuable therapeutic proteins. Ever since we demonstrated expression of human-elastin derived polymers for various biomedical applications (Guda et al., 2000), we have extended this approach to express vaccines antigens for Cholera, Anthrax (Daniell et al., 2001, Daniell 2003), monoclonal antibody (Daniell et al., 2001) and human therapeutic proteins, including Human Serum Albumin, (Fernandez et al., 2003), Magainin (DeGray et al., 2001), Interferon (Daniell 2003) and Insulin like Growth Factor (Daniell, 2003). Several other laboratories have expressed Human Somatotropin Interferon-GUS fusion proteins to improve stability (Reddy et al., 2003) and tetanus vaccine antigens (Tregoning et al., 2003) in transgenic chloroplasts. Without any exception, all of these therapeutic proteins have been expressed in transgenic tobacco

chloroplasts, meeting the zero-tolerance of food crops for plant-derived pharmaceuticals advocated by various environmental groups.

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However, there is an urgent need for oral delivery of therapeutic proteins and vaccine antigens to dramatically reduce their production, purification, storage and transportation costs and minimize complications associated with intravenous delivery. Carrot (Daucus carota L.) is one of the most important vegetables used worldwide for human and animal consumption, due to its excellent source of sugars, vitamins A, C and fiber in the diet. The carrot plant is biennial, completing its life cycle in two years. In the first year the plant produces the fleshy taproot, which is edible. If left in the ground, plants flower in the second year after passing through a cold season (Yan, W. & Hunt, L.A Reanalysis of Vernalization Data of Wheat and Carrot, Annals of Botany 84, 615-619 (1999). In addition, chloroplast genomes in the cultivated carrot crop are transmitted strictly through maternal inheritance (Vivek et al 1999). Thus, carrot is environmentally safe and is doubly protected against transgene flow via pollen and seeds to achieve zero-tolerance on transgene flow advocated for food crops. Carrot somatic embryos are single cell derived and multiply through recurrent embryogenesis; this provides uniform source of cell culture, which is one of the essential requirements for producing therapeutic proteins (homogeneous single source of origin). Carrot cells divide rapidly and large biomass is produced using bioreactors. Cultured carrot cells are edible and could be used directly to deliver precise dose of vaccine antigens or biopharmaceuticals. When delivered via edible carrots, there is no need to cook and this would preserve the structural integrity of therapeutic proteins during consumption. Viable for long duration on culture medium, encapsulated embryos are used as synthetic seeds for cryopreservation and controlled germination (Tessereau, H., B. Florin, M. C. Meschine, C. Thierry and V. Pétiard, 1994). Thus, transgenic carrot with enhanced medicinal or nutritional value can play a vital role in improving human or animal health

However, in order to engineer the carrot chloroplast genome, one has to overcome several major hurdles. So far, the chloroplast genome has been transformed only by using green leaves as explants, which contain large chloroplasts. The first challenge is to introduce foreign DNA into small proplastids and identify appropriate regulatory sequences and selectable markers that function in non-green plastids. The

second challenge is to regenerate chloroplast transgenic plants via somatic embryogeneis and achieve homoplasmy, which lacks the benefit of subsequent rounds of selection offered by organogenesis, while using leaves as explants. Because of these reasons, chloroplast genetic engineering has been achieved only in a few solanaceous crops other than tobacco, such as tomato (Ruf et al., 2001) and potato (Sidorov et al., 1999), although the later crop remained sterile. Non-solanaceous crops continue to be a challenge to transform, even though regeneration was possible from green leaves via organogenesis. For example, *Arabidopsis* transgenic plants were sterile (Sidkar et al., 1998) and even stable integration or homoplasmy could not be achieved in oilseed rape (Bing Kai Hou et al., 2003).

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Table 1 shows an exemplary list of the development of transgene expression in chloroplasts.

TABLE 1

# Transgene Expression in chloroplasts

Agronomic traits	Gene	Promoter	5'/3' Regulatory elements	Reference
Insect resistance	Cry1A(c)	Prrn	rbcL / Trps16	Mc Bride et al
			×.	1995
Herbicide	CP4 (petunia)	Prrn	ggagg / TpsbA	Daniell et al
resistance				1998
Insect resistance	Cry2Aa2	Prrn	ggagg (native) / TpsbA	Kota et al 1999
Herbicide	CP4 (bacterial or	Prrn	rbcL or T7 gene 10 / Trps16	Ye at al 2001
resistance	synthetic)			
Insect resistance	Cry2Aa2 operon	Prrn	Native 5'UTRs / TpsbA	DeCosa et al
·				2001
Disease resistance	MSI-99	Prrn	ggagg / TpsbA	DeGray et al
				2001
Salt and drought	tps	Prrn	ggagg / TpsbA	Lee et al 2003
tolerance				
Phytoremediation	merA <sup>a</sup> /merB <sup>b</sup>	Prrn	ggagg <sup>a, b</sup> / T <i>psb</i> A	Ruiz et al 2003

Biopharmaceutical	Gene	Promoter	5'/3'	% tsp	Reference
proteins			regulatory	expression	
			elements		
Protein based polymer	EG121	Prrn ,	T7gene10 /	Not tested	Guda et al
		,	TpsbA		2000
Human somatotropin	hST	Prrn <sup>a</sup> ,	T7gene10 <sup>a</sup> or	7.0 % and	Staub et al
		$PpsbA^b$	psbA <sup>b</sup> / Trps16	1.0% <sup>b</sup>	2000
Cholera toxin	ctxB	Prrn	ggagg / T <i>psb</i> A	4%	Daniell et al
					2002
Tetanus toxin	TetC	Prrn	T7 gene 10 <sup>a</sup> ,	25% a, 10% b	Tregoning et al
	(bacterial		atpBb / TrbcL		2003
	and				
	synthetic)				
Human Serum Albumin	hsa	Prrna,	ggagg <sup>a</sup> , psbA <sup>b</sup> /	0.02% a,	Fernandez-San
		PpsbA <sup>b</sup>	TpsbA	11.1% <sup>b</sup>	Milan et al
					2003
Interferon alpha 5	INF□5	Prrn	PpsbA/TpsbA	ND	Torres
Interferon alpha 2B	INF□2B	Prrn	PpsbA/TpsbA	19%	· Falconer

Table 1(continued)

ifn-g	P <i>psb</i> A	PpsbA/TpsbA	6%	Leelavathi and
				Reddy, 2003
	Prrn	ggagg / TpsbA	ND	Daniell et al
				(photosynthesi
				s)
Igf-1	Prrn	PpsbA/TpsbA	33%	Ruiz G
		·		
Pag	Prrn	PpsbA/TpsbA	4-5%	Watson
CaF1~Lcr	Prrn	PpsbA/TpsbA	4.6 %	Singleton
V				
	Igf-1 Pag CaF1~Lcr	Prrn	Prrn         ggagg / TpsbA           Igf-1         Prrn         PpsbA/TpsbA           Pag         Prrn         PpsbA/TpsbA           CaF1~Lcr         Prrn         PpsbA/TpsbA	Prrn         ggagg / TpsbA         ND           Igf-1         Prrn         PpsbA/TpsbA         33%           Pag         Prrn         PpsbA/TpsbA         4-5%           CaF1~Lcr         Prrn         PpsbA/TpsbA         4.6 %

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## SUMMARY OF THE INVENTION

One aspect of this invention describes methods for transforming plastids using a highly efficient process for carrot plastid transformation through somatic embryogenesis. Still other aspects of this invention provide for vectors which are capable plastid transformation through somatic embryogenesis. Still another aspect provides for transformed plastids, plants, and plant parts, which have been transformed through somatic embryogenesis through the methods and vectors described herein. This application, along with the knowledge of the art, provides the necessary guidance and instructions to engineer the plastid genome of several major crops in which regeneration is mediated through somatic embryogenesis. Cereal crops (wheat, rice, corn, sugarcane), legumes (soybean, alfalfa), oil crops (sunflower, olive), cash crops (cotton, coffee, tea, rubber, flex, cork oak, pines), vegetables (eggplant, cucumber, cassava, chili pepper, asparagus etc.), fruits (apple, cherry, banana, plantain, melons, grape, guava), nuts (cashew, walnuts, peanuts), and trees (date palm etc.) are regenerated through somatic embryogenesis. Another aspect of this invention shows the constructs of chloroplast vectors for a variety of different species using the same primers( universal plastid primers).

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## BRIEF DESCRIPTION OF THE FIGURES

Fig. 1 (A-B). shows the physical map of the carrot chloroplast transformation vectors.

Fig 1(A) shows carrot chloroplast transformation vector pDD-Dc-gfp/BADH carries the gfp and BADH genes expressed under the regulation of T7 gene 10 5' untranslated region (UTR)/ rps16 3'UTR and PpsbA 5' and 3' UTR respectively. The Prrn promoter of 16S r-RNA gene, having both PEP and NEP recognition sites, drives expression of the cassette.

Fig 1(B) shows the carrot chloroplast transformation vector pDD-Dc-aadA/BADH carries the aadA and BADH gene. Expression of aadA gene is under the regulation of Shine-Dalgarno sequence and psbA 3'UTR while that of BADH is regulated by gene10 5' and rps16 3'UTR. AfIIII/PvuII digested  $\sim$  4.9 kb DNA fragment used as a probe for Southern analysis of the transgenic plants and landing

sites for primers 3P/3M and 16SF/1M used to confirm the presence of the transgene integration into carrot plastids are shown.

Fig. 2(A-C) shows the expression of GFP in carrot cultures transformed with chloroplast vector pDD-*Dc-gfp/BADH*; visible under confocal fluorescent microscope at fluorescence emission in green at 488nm blue Argon (laser).

Fig. 2(A) shows the untransformed control carrot culture,

Fig. 2(B) shows the transformed embryogenic calli,

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Fig.2(C) shows the transformed embryogenic carrot calli differentiated into globular somatic embryos and

Fig. 2(D) shows a somatic embryo differentiated into cotyledonary stage.

Fig. 3 (A-D) shows the visual selection of green transgenic cells versus yellow non-transgenic carrot cells culture.

Fig. 3(A-B) shows the transgenic carrot cell culture turned green due to the expression BADH (Plate A) while wild type culture remained yellow (Plate B). Transgenic carrot cell culture can be distinguished as green-transgenic cell culture vs yellow non-transgenic carrot cell culture, when heteroplasmic transgenic cell line was placed on medium without any selection agent (Plate C and D).

Fig. 4 (A-C) shows the transgene (aadA and badh) integration into the carrot plastid genome was confirmed by PCR and Southern blot analysis.

Fig 4(A) shows the use of internal primers 3P (land on flanking sequence) and 3M (land on *aadA* gene) ~1.65 kb size PCR product was amplified at 64°C annealing temperature, confirmed transgene integration into plant cell lines.

Fig. 4(B) shows the use of a set of primer 16SF (landing on the native chloroplast genome) and 1M (landing on the *aadA* gene) yield ~ 2.5 kb size PCR product at 64°C annealing temperature, confirmed plastid specific integration of the transgenes. Lane 2 stand for DNA from non-transgenic carrot cells and lanes 2-9 represents DNA from seven transgenic carrot cell lines. Primers landing sites for primer pairs (3P/3M and 16SF/1M) is shown in Fig. 1B.

Fig 4(C) shows the southern blot analysis of plastid genome of untransformed and transformed carrot with vector pDD-*Dc-aadA/BADH*. Carrot genomic DNA (5 μg per lane) digested with AfIIII and PvuII and transferred to nitrocellulose membrane, was hybridized with the 4.9 kb radioactive labeled P<sup>32</sup> DNA probe (containing 2.4 kb

flanking sequence and 2.5 kb transgene sequence, see Fig. 1B). Lane 1, control DNA from untransformed transgenic plant showed 2.4 kb size band while heteroplasmic transgenic plant from cell line one, lane 2 showed both bands. Homoplasmy in transgenics plants from different cell lines (lanes 3-8) was achieved by repetitive subcultures of transgenic cells in liquid medium.

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Fig. 5 (A-B) shows BADH enzyme activity (nmol/min/mg/protein) and BADH expression was analyzed in protein extracts from untransformed and transformed carrot with plastid vector pDD-*Dc-aadA/BADH*.

Fig 5(A) shows the reduction of NAD<sup>+</sup> dependent BADH enzyme was analyzed for the formation of NADH at 340 nm in presence of betaine aldehyde. Very low BADH activity was detected in untransformed cells suspension (U), carrot root (U) and leaf (U). On the other hand, high BADH activity was recorded about 54 % in transformed carrot cells suspension (T), about 72% in root (T) in comparison to leaf (T) tissues.

Fig. 5(B) shows that the BADH expression was analyzed by western blot. Whole cell extracts from transformed and untransformed carrot cell culture, root and leaf tissues were prepared and 50 μg total soluble protein from each sample was run on 10% SDS-PAGE and protein transferred to Immuno-blot<sup>TM</sup> PVDF membrane and hybridized with polyclonal anti-BADH serum, raised in rabbits against native BADH. Antigenic peptides were detected using horseradish peroxidase-linked secondary antibody. Lane 1, 2, 3 contain whole carrot extract from untransformed cell culture, root and leaf and lane 4, 5, 6 contain whole carrot extract from transformed cell culture, root and leaf. Transgenic cells expresses about 50 % less (lane 4), root about 25 % less (lane 5) BADH protein than leaf (lane 6).

Fig. 6 (A-C) shows the effect of different salt concentrations on growth of untransformed and transformed carrot cell suspension cultures with chloroplast vector pDD-Dc-aadA/BADH.

Fig. 6(A) shows the dry mass in untransformed carrot cell culture.

Fig. 6(B) shows transformed cell cultures produced in liquid medium containing 100 mM NaCl.

Fig. 6(C) shows the stimulation of BADH activity in presence of salt. Untransformed and transformed carrot cells in suspension cultures were placed on

shaker at 130 rpm speed for two weeks in liquid medium containing 0, 100, 200 and 300 mM NaCl. Elevated level of BADH activity in transgenic cell cultures was noticed when liquid growth medium containing 100 and 200 mM NaCl.

Fig. 7 Effect of salt (100–500 mM NaCl) on untransformed (U) and transformed (T) carrot plants. Transgenic plants were tested for one month on different concentration of NaCl. Plants were irrigated with water containing different concentrations NaCl at alternative days up to four weeks.

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Fig. 8 shows the plasmid pDD-Ta-aphA-6/nptII. More particularly the plasmid illustrates pDA-76 (aphA-6/nptII expression cassette), having a backbone vector pBluescript II KS, a selectable marker/host cell Ampicillin/Kan/ XL-1 Blue MRF' Tc, and a flanking region from Triticum aestivum (Ta).

Fig 9 shows the plasmid pDD-So-aphA-6/nptII. More particularly the plasmid illustrates pDA-76 (aphA-6/nptII expression cassette), having a backbone vector pBluescript II KS, a selectable marker/host cell Ampicillin/Kan/ XL-1 Blue MRF' Tc, and a flanking region from Saccharum officinarum (So).

Fig 10 shows the plasmid pDD-Dc-aphA-6/nptII. More particularly the plasmid illustrates pDA-76 (aphA-6/nptII expression cassette), having a backbone vector pBluescript II KS, a selectable marker/host cell Ampicillin/Kan/ XL-1 Blue MRF' Tc, and a flanking region from *Daucus carota* (Dc).

Fig 11 shows the plasmid pDD-Dc-aadA/BADH. More particularly the plasmid illustrates pDA-29 (aadA/BADH expression cassette) having a backbone vector pBluescript II KS, a selectable marker/host cell Ampicillin/spectinomycinXL-1 Blue MRF' Tc, and a flanking region from *Daucus carota* (Dc).

Fig. 12 shows the plasmid pDD-Dc-gfp/BADH. More particularly the plasmid pDA-30 (gfp/BADH expression cassette), having a backbone vector pBluescript II KS, a selectable marker/host cell Ampicillin/Kan/ XL-1 Blue MRF' Tc, and a flanking region from *Daucus carota* (Dc).

Fig. 13 shows the plasmid pDD-Gh-aphA-6/nptII. More particularly the plasmid illustrates pDA-76 (aphA-6/nptII expression cassette), having a backbone vector pBluescript II KS, a selectable marker/host cell Ampicillin/Kan/ XL-1 Blue MRF' Tc, and a flanking region from Gossypium hirsutum (Gh).

Fig. 14 shows the plasmid pDD-Gh-aadA/BADH. More particularly the plasmid illustrates pDA-29 (aadA/BADH expression cassette), having a backbone vector pBluescript II KS, a selectable marker/host cell Ampicillin/spectinomycinXL-1 Blue MRF' Tc and a flanking region from Gossypium hirsutum (Gh).

Fig. 15 shows the plasmid pDD-Gh-gfp/BADH. More particularly the plasmid illustrates pDA-30 (gfp/BADH expression cassette), having a backbone vector pBluescript II KS, a selectable marker/host cell Ampicillin/XL-1 Blue MRF' Tc, and a flanking region from Gossypium hirsutum (Gh).

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Fig. 16 shows the plasmid pDD-Zm-aadA/BADH. More particularly the plasmid illustrates pDA-29 (aadA/BADH expression cassette), having a backbone vector pBluescript II KS, a selectable marker/host cell Ampicillin/spectinomycinXL-1 Blue MRF' Tc, and a flanking region from Zea mays (Zm).

Fig. 17 shows the plasmid pDD-Zm-gfp/BADH. More particularly the plasmid illustrates pDA-30 (gfp/BADH expression cassette), having a backbone vector pBluescript II KS, a selectable marker/host cell Ampicillin/XL-1 Blue MRF' Tc, and a flanking region from Zea mays (Zm).

Fig. 18 shows the plasmid pDD-Zm-aphA-6/nptII. More particularly the plasmid illustrates pDA-76 (aphA-6/nptII expression cassette), having a backbone vector pBluescript II KS, a selectable marker/host cell Ampicillin/Kan/ XL-1 Blue MRF' Tc, and a flanking region from *Zea mays* (Zm).

Fig. 19 shows the plasmid pDD-Pv-aphA-6/nptII (switchgrass). More particularly the plasmid illustrates pDA-76 (aphA-6/nptII expression cassette), having a backbone vector pBluescript II KS, a selectable marker/host cell Ampicillin/Kan/XL-1 Blue MRF' Tc, and a flanking region from *Panicum virgatum* (Pv).

Fig. 20 shows the plasmid pDD-Pv-aadA/BADH (switchgrass). More particularly the plasmid illustrates pDA-29 (aadA/BADH expression cassette), having a backbone vector pBluescript II KS, a selectable marker/host cell: Ampicillin/spectinomycinXL-1 Blue MRF' Tc, and a flanking region from *Panicum virgatum* (Pv).

Fig.21 shows the plasmid pDD-Cd-aphA-6/nptII (bermudagrass). More particularly the plasmid illustrates pDA-76 (aphA-6/nptII expression cassette), having

a backbone vector pBluescript II KS, a selectable marker/host cell: Ampicillin/Kan/XL-1 Blue MRF' Tc, and a flanking region from *Cynodon dactylon* (Cd).

Fig. 22 shows the plasmid pDD-Nt-aphA-6/nptII. More particularly the plasmid illustrates pDA-76 (aphA-6/nptII expression cassette), having a backbone vector pBluescript II KS, a selectable marker/host cell Ampicillin/Kan/ XL-1 Blue MRF' Tc, and a flanking region from *Nicotiana tabacum* (Nt).

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Fig. 23 shows the plasmid pDD-Os-aphA-6/nptII. More particularly the plasmid illustrates pDA-76 (aphA-6/nptII expression cassette), having a backbone vector pBluescript II KS, a selectable marker/host cell Ampicillin/Kan/ XL-1 Blue MRF' Tc, and a flanking region from *Oryza sativa* (Os).

Fig. 24 shows the plasmid pDA-66. More particularly the plasmid illustrates psbA 5'UTR BACKBONE VECTOR pUC 19, which is a Derivative of pLD-CtV basic vector (modified MCS) having a selectable marker/host cell Ampicillin/Kan/ XL-1 Blue MRF' Tc, and a flanking region from Tobacco.

Fig. 25 shows the plasmid pDD-Ta-aadA/BADH. More particularly the plasmid illustrates pDA-29 (aadA/BADH expression cassette), having a backbone vector pBluescript II KS, a selectable marker/host cell Ampicillin/spectinomycinXL-1 Blue MRF' Tc, and a flanking region from *Triticum aestivum (Ta)*.

Fig. 26 shows the plasmid pDD-Ta-gfp/BADH. More particularly the plasmid illustrates pDA-30 (gfp/BADH expression cassette), having a backbone vector pBluescript II KS, a selectable marker/host cell Ampicillin/XL-1 Blue MRF' Tc, and a flanking region from *Triticum aestivum* (Ta).

Fig. 27 shows the plasmid pDD-Hv-aphA-6/nptII. More particularly the plasmid illustrates pDA-76 (aphA-6/nptII expression cassette), having a backbone vector pBluescript II KS, a selectable marker/host cell Ampicillin/Kan/ XL-1 Blue MRF' Tc, and a flanking region from *Hordeum vulgare (Hv)*.

Fig. 28 is a schematic view of a Double Barreled Plastid Vector harboring aphA-6 and aphA-2 genes conferring resistance to aminoglycosides according to the description contained herein.

Fig. 29A and 30A illustrate the construction of maize chloroplast transformation vector, where flanking regions were amplified using PCR. The PCR products were cloned and the expression cassette was inserted in the transcriptionally

active spacer region between trnI/trnA genes. The expression cassette of Fig. 30A has the Prrn promoter driving the expression of GFP and BADH, which are regulated by (5') gene10/rps16 3' and psbA 5'/3' UTRs respectively. The expression cassette of 31A has the Prrn promoter driving the expression of aadA and BADH. The latter gene is regulated by (5') gene10/rps16 3' UTRs.

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Figs. 29B and 30B shows the functions of the genes in the maize chloroplast transformation vectors which were tested in E. coli. For observing GFP expression, cells were plated on LB agar (Amp) plates and incubated at 37°C overnight. Cells harboring pDD34-ZM-GFP-BADH were seen to fluoresce when exposed to UV light, as is seen in Fig 30B. To test the aadA gene expression, cells harboring pDD33-ZM-aadA-BADH plasmid were plated on LB agar plates containing spectinomycin (100mg/ml) and incubated at 37°C overnight. Transformed cells grow on spectinomycin, as can be seen in Fig. 31B.

Fig. 31 shows GFP expression in embryogenic maize cultures studied under the confocal microscope. Fig. 32A is a non-transgenic control, while Figs. 32B-C are transformed maize embryogenic calli. The selection in Figs. 30-31 was initiated two days after bombardment by transferring the bombarded calli to callus induction medium containing BA or streptomycin. After eight weeks, a number of the healthy growing calli from different bombardment experiments were examined for GFP expression under the fluorescent stereomicroscopeand the confocal microscope. Somatic embryos were regenerated on maize regeneration medium containing BA or streptomycin.

Figs. 32(A-B) shows maize plants on regeneration medium containing streptomycin or betaine aldehyde. Fig. 32A illustrates maize chloroplast transgenic plants which were capable of growth on the selection agent indicating that construction of transgenic maize, while untransformed maize plants did not grow on the selection medium.

Fig 32B shows PCR confirmation of chloroplast transgenic plants using appropriate primers. Lanes 1-3, plants transformed with pDD34-ZM-gfp-BADH and Lanes 4-5, plants transformed with pDD33-ZM-aadA-BADH. Lanes – and + represent the negative and positive controls respectively. Genomic DNA was isolated from the

leaf tissues and PCR was performed on transformed and non-transformed tissues using appropriate primers.

Fig. 33(A-C) shows the Transformed cotton cultures (Gossypium hirsutum cv. Coker310FR) with chloroplast vector pDD-C-aphA6/aphA2; selected on medium MST1 (0.1 mg/l 2,4-D and 0.5 mg/l kinetin) supplemented with 50 mg/l kanamycin.

Fig. 33(A) shows the untransformed control cotton calli.

Fig. 33(B) shows the transformed primary cotton calli,

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Fig. 33(C) shows the transformed cotton calli subcultured from transgenic primary cotton calli (Plate B).

Fig 34 (A-B) show the transgene (aphA6 and aphA2) integration into the cotton plastid genome was confirmed by PCR.

Fig. 34(A) shows the use of internal primers 3P (land on flanking sequence) and aphA6-rev (land on aphA6 gene) ~1.7 kb size PCR product was amplified at 64°C annealing temperature, confirmed transgene integration into cotton calli.

Fig 34(B) shows the use of a set of primer 16SF (landing on the native chloroplast genome) and aphA6-rev (landing on the aphA6 gene) yield ~ 2.5 kb size PCR product at 64°C annealing temperature, confirmed plastid specific integration of the transgenes. Lane 1 represents the 1kb Plus molecular marker (ladder). Lane 2 stand for DNA from non-transgenic cotton calli and lane 3 represents DNA from transgenic cotton calli selected on 50 mg/l kanamycin.

Fig. 35 shows the sequence of the aadA/BADH expression cassette (SEQ ID No. 1).

Fig. 36 shows the sequence of the gfp/BADH expression cassette (SEQ ID No. 2).

Fig. 37 shows the sequence of the aphA-6/nptII expression cassette (SEQ ID No. 3).

Fig. 38 shows a schematic view of a general plastid transformation vector.

## DETAILED DESCRIPTION OF THE INVENTION

In one aspect of the invention homoplasmic plants regenerated from the plant cell cell culture via somatic embryogenesis is provided. Another aspect of this invention is plastid transformation vectors capable for use in non-green explants which

can lead to somatic embryogenesis of the plant cells. Yet another aspect of this invention provides for transgene expression in non-green edible plant parts. Aspects of this invention further describe transformation of monocots, legumes, vegtables, fruit crops, and transgene expression within the non-green plant parts of these plants. Another aspect of this invention provides for the expression of heterologous proteins using a plastid transofrmation vector suitable for transforming the non-green plant parts. In other aspects methods of transforming plastid genomes to express, via somatic embryogenesis, heterologous proteins is provided transformed plants and progeny thereof, which express the protein of interest. Yet another aspect of this invention is the introdcution of foreign DNA into the small proplastids of plants, and the identification of selectable markers which function in non-green plastids. Still other aspects of this invention provide for regenerated chloroplast transgenic plants via somatic embryogenesis to achieve homoplasmy.

The preferred aspects of this application are applicable to all plastids of higher plants. These plastids include the chromoplasts, which are present in the fruits, vegetables, and flowers; amyloplasts which are present in tubers such as potato; proplastids in the roots of higher plants; leucoplasts and etioplasts (which express in the dark), both of which are present in the non-green parts of plants. The aspects of this application are also applicable to various developmental stages of chloroplast, wherein the chloroplast are not fully green.

## **Definitions**

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To better understand the current disclosure, the following definitions, which shall hold their meaning throughout this application unless otherwise noted, are provided to put the application in proper context.

Heterologous generally means derived from a separate genetic source. Of course this invention contemplates the use of heterologous and homologous DNA, as well as operons suitable for expression in plant plastids.

An expression cassette, is generally understood in the art as a cloning vector that contains the necessary regulatory sequences to allow transcription and translation of a cloned gene or genes.

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Properly folded should be understood to mean a protein that is folded into its normal conformational configuration, which is consistent with how the protein folds as a naturally occurring protein expressed in its native host cell.

When refering to plants throughout the application it should be understood that the current aspects disclosed herein contemplate the transformation of the plastids of all organisms and plants which contain plastids. For purposes of clarity the phrase "higher plants" generally includes solanaceous and non-solanaceous plants, and the exemplary list of crops, fruits, flowers, vegtables, beans, medicinal plants, and all other plants which one skilled in the art would recognize as being a higher plant.

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Substantially homologous as used throughout the ensuing specification and claims, is meant a degree of homology to the native Human Serum Albumin sequence in excess of 50%, most preferably in excess of 80%, and even more preferably in excess of 90%, 95% or 99%. Substantial sequence identity or substantial homology as used herein, is used to indicate that a nucleotide sequence or an amino acid sequence exhibits substantial structural or functional equivalence with another nucleotide or amino acid sequence. Any structural or functional differences between sequences having substantial sequence identity or substantial homology will be de minimis; that is, they will not affect the ability of the sequence to function as indicated in the desired application. Differences may be due to inherent variations in codon usage among different species, for example. Structural differences are considered de minimis if there is a significant amount of sequence overlap or similarity between two or more different sequences or if the different sequences exhibit similar physical characteristics even if the sequences differ in length or structure. Such characteristics include, for example, ability to maintain expression and properly fold into the proteins conformational native state, hybridize under defined conditions, or demonstrate a well defined immunological cross-reactivity, similar biopharmaceutical activity, etc. Each of these characteristics can readily be determined by the skilled practitioner in the art using known methods.

Non-green plastids generally refers to any plastid that is not green. Examples of such plastids include, the chromoplasts, which are present in the fruits, vegetables, and flowers; amyloplasts which are present in tubers such as potato; proplastids in the roots of higher plants; leucoplasts and etioplasts (which express in the dark) and

different develop stages of chloroplast, wherein the chloroplast is not green. Further the non-green part of plants and plant cells is well characterized and understood in the art.

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Spacer region is understood in the art to be the region between two genes. The chloroplast genome of plants contains spacer regions which highly conserved nuclear tide sequences. The highly conserved nature of the nuclear tide sequences of these spacer regions chloroplast genome makes the spacer region ideal for construction of vectors to transform chloroplast of a wide variety of plant species, without the necessity of constructing individual vectors for different plants or individual crop species. It is well understood in the art that the sequences flanking functional genes are well-known to be called "spacer regions". The special features of the spacer region are clearly described in the Applicant's Application No. 09/079,640 with a filing date of May 15, 1998 and entitled UNIVERSAL CHLOROPLAST INTEGRATION OF EXPRESSION VECTORS, TRANSFORMED PLANTS AND **PRODUCTS** THEREOF. The aforementioned Application No. 09/079,640 is hereby incorporated by reference. It was well-known that there are at least sixty transcriptionally-active spacer regions within the higher plant chloroplast genomes (Sugita, M., Sugiura, M., Regulation of Gene Expression in Chloroplasts of Higher Plants, Plant Mol. Biol., 32: 315-326, 1996). Specifically, Sugita et al. reported sixty transcriptionally-active spacer regions referred to as transcription units, as can be seen in Table II of the article. Because the transcriptionally active spacer regions are known, a universal vector, as described in the Applicant's U.S. Patent Application No. 09/079,640, can be used in the identified spacer regions contained within a variety of the higher plant chloroplast genomes. By utilizing the teachings in Sugita et al., intergenic spacer regions are easily located in the plastid genome. Consequently this allows one skilled in the art to use the methods taught in the Applicant's U.S. Patent Application No. 09/079,640 to insert a universal vector containing the psbA, the 5' untranslated region (UTR) of psbA and the gene coding for the protein of interest into the spacer regions identified by Sugita et al., and found across higher plants. The aforementioned applications and article are incorporated by reference.

Selectable marker provides a means of selecting the desired plant cells, vectors for plastid transformation typically contain a construct which provides for expression

of a selectable marker gene. Marker genes are plant-expressible DNA sequences which express a polypeptide which resists a natural inhibition by, attenuates, or inactivates a selective substance, i.e., antibiotic, herbicide, or an aldehyde dehydrogenase such as Betaine aldehyde dehydrogenase (described in the Applicant's Application No. 09/807,722 filed on April 18, 2001, and herein fully incorporated by reference). Alternatively, a selectable marker gene may provide some other visibly reactive response, i.e., may cause a distinctive appearance or growth pattern relative to plants or plant cells not expressing the selectable marker gene in the presence of some substance, either as applied directly to the plant or plant cells or as present in the plant or plant cell growth media.

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In either case, the plants or plant cells containing such selectable marker genes will have a distinctive phenotype for purposes of identification, i.e., they will be distinguishable from non-transformed cells. The characteristic phenotype allows the identification of cells, cell groups, tissues, organs, plant parts or whole plants containing the construct. Detection of the marker phenotype makes possible the selection of cells having a second gene to which the marker gene has been linked.

The use of such a marker for identification of plant cells containing a plastid construct has been described in the literature. In the examples provided below, a bacterial aadA gene is expressed as the marker. Expression of the aadA gene confers resistance to spectinomycin and streptomycin, and thus allows for the identification of plant cells expressing this marker. The aadA gene product allows for continued growth and greening of cells whose chloroplasts comprise the selectable marker gene product. Numerous additional promoter regions may also be used to drive expression of the selectable marker gene, including various plastid promoters and bacterial promoters which have been shown to function in plant plastids.

Inverted Repeat Regions are regions of homology, which are present in the inverted repeat regions of the plastid genome (known as IRA and IRB), two copies of the transgene are expected per transformed plastid. Where the regions of homology are present outside the inverted repeat regions of the plastid genome, one copy of the transgene is expected per transformed plastid.

Structural equivalent should generally be understood meaning a protein maintaining the conformational structure as the native protein expressed in its natural cell.

Vectors

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The current application contemplates the use of vectors capable of plastid transformation, particularly for plastid transformation. Such vectors include plastid expression vectors such as pUC, pBR322, pBLUESCRIPT, pGEM, and all others identified by Daniel in U.S. Patent No. 5,693,507 and U.S. Patent No. 5,932,479. Included are also vectors whose flanking sequences are located outside of the embroidered repeat of the chloroplast genome. These publications and patents are hereby incorporated by reference to the same extent as if each individual publication or patent was specifically an individually indicated to be incorporated by reference.

The universal vector is described in WO 99/10513 which was published on March 4, 1999, and Application No. 09/079,640 which was filed on May 15, 1998, wherein both of said references are incorporated in their entirety.

Basic pLD vector, developed in this laboratory for chloroplast transformation, was used (Daniell *et al.*, 1998; Daniell *et al.*, 2001b; De Cosa *et al.*, 2001; Guda *et al.*, 2000; Kota *et al.*, 1999). High levels of foreign protein expression in chloroplasts (3–21% of tsp) have been shown for different proteins using the SD 5' sequence (Daniell et al., 2001b; DeGray et al., 2001; Kota et al., 1999).

It should be noted that the vectors described herein are illustrative examples and vectors can be constructed with different promoters such as was described in U.S. Patent Application No. 09/079,640, different selectable markers such as those described in U.S. Patent Application No. 09/807,722, and different flanking sequences suitable for integration into a variety of plant plastid genomes.

## GENERAL METHODOLGY FOR TRANSFORMING THE PLASTID GENOME

This illustrative example shows generally all of the necessary steps to practice the Applicants invention. Of course other suitable methods, which are known in the art may be substituted or used to supplement the example methodology described herein.

## Isolation of genomic DNA from plants.

Mortar and pestle, liquid nitrogen, fresh dark green leaves. DNeasy Plant Mini Kit (QIAGEN Inc.)

## PCR amplification of chloroplast flanking sequence.

Materials for PCR reaction: Genomic DNA (50-100ng/μl), dNTPs, 10x pfu buffer, Forward primer, Reverse primer, autoclaved distilled H<sub>2</sub>O and Turbo pfu DNA Polymerase.

## 5 Vector construction.

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- 1. Plasmid pUC19 or pBlueScript SK (+/-).
- 2. Species specific PCR amplified chloroplast DNA flanking sequences.
- 3. A promoter functional in plastids, 5'UTR of chloroplast gene, selectable marker gene, gene of interest and chloroplast 3'UTR.
  - 4. Restriction enzymes and buffers.
- 5. T4 DNA polymerase to remove 3' overhangs to form blunt ends and fill-in of 5' overhangs to form blunt ends or Klenow large fragment (fill-in of 5' overhangs to form blunt ends), alkaline phoshatase for dephoshorylation of cohesive ends, DNA ligase to form phosphodiester bonds and appropriate buffers.
  - 6. Water baths or incubators set at different temperatures.

## Preparation for biolistics.

- 1. Autoclaved Whatman filter paper #1 (55 mm in diameter) dried in oven.
- 2. 100% ethanol.
- 3. Autoclaved tips in box, autoclaved kimwipes tissues wrapped in aluminum foil.
  - 4. Sterile gold particles stored at -20°C in 50% glycerol (see Notes 1 and 2).
  - 5. Sterile rupture discs (1100 psi) and macrocarriers sterilized by dipping in 100% ethanol.
    - 6. Autoclaved steel macrocarrier holders and stopping screens.
  - 7. Freshly prepared 2.5 mM CaCl<sub>2</sub>: weigh 1.84 g and dissolve in 5 mL  $H_2O$  and filter sterilized with 0.2  $\mu$ m filter.
  - 8. 0.1 M spermidine (highly hygroscopic): dilute 1M spermidine stock to 10x and aliquot  $100 \mu L$  in 1.5 mL Eppendrop tubes to store at -20°C. Discard each tube after single use.

#### Medium preparation for plant tissue culture.

#### 2.5.1. Tobacco.

Medium for 1000 mL: 4.3 g MS salts (INVITROGEN Inc.), H<sub>2</sub>O (molecular biology grade), 100 mg/L myo-inositol, 1 mg/L thiamine-HCl, 3% sucrose for shoot induction and 2% sucrose for root induction, 1mg/L 6-benzyl aminopurine (BAP; use 1 mL from 1mg/mL stock), 0.1 mg/L indole-3- acetic acid (use 0.1 mL from 1 mg/mL IAA stock), 1 mg/L indole-3-butyric acid for root induction (use 1 mL from 1mg/mL IBA stock). Add 500 mg/L spectinomycin in autoclaved medium when it cools to 45°C - 50°C (use 5 mL filter sterilized spectinomycin from 100 mg/mL stock).

Edible crops.

Potato.

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Medium for 1000 mL: 4.3 g MS salts, B5 vitamins (make 100x solution in 100 mL H<sub>2</sub>O by dissolving: 1 g myo-inositol, 10 mg nictonic acid, 10 mg pyridoxine-HCl, 100 mg thiamine-HCl; use 10 mL, store remaining solution at 4°C), 5 mg/l zeatin riboside (use 0.5 mL from 1 mg/mL ZR stock), 0.1 mg/l α-napthaleneacetic acid (use 0.1 mL from 1 mg/mL NAA stock), 40 to 500 mg/L spectinomycin.

**Tomato** 

Medium for 1000 mL: 4.3 g MS salts, B5 vitamins (10 mL from 10x stock), 0.2 mg/l indole-3-acetic acid (use 0.2 mL from 1 mg/mL IAA stock), 3 mg/l of 6-benzylaminopurine (use 3 mL from 1 mg/mL BAP stock). 300 or 500 mg/L spectinomycin.

For all plant growth media adjust to pH 5.8 with 1N KOH or 1N NaOH and add 6g/L phytagel (Sigma) before autoclaving at 121°C for 20 min. For preparation of 1mg/mL stock of BAP, IAA, IBA, NAA, ZR respectively: weigh 10 mg powder and dissolve first in 1 or 2 drops of 1N NaOH and make up the final volume to 10 mL; store all plant growth regulators at 4°C for 1-2 months).

## Molecular analysis of transgenic plants.

PCR analysis for gene integration into tobacco chloroplasts

PCR reaction for 50  $\mu$ L: 1.0  $\mu$ l genomic DNA (50-100 ng/ $\mu$ l), 1.5  $\mu$ l dNTPs (stock10 mM), 5.0  $\mu$ l (10x PCR buffer), 1.5  $\mu$ l Forward primer (to land on the native chloroplast genome; stock 10  $\mu$ M), 1.5  $\mu$ l Reverse primer (to land on the transgene; stock 10  $\mu$ M), 39.0  $\mu$ l autoclaved distilled H<sub>2</sub>O and 0.5  $\mu$ l Taq DNA polymerase.

Analysis of homoplasmy by Southern blots.

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1. Depurination solution: 0.25 N HCl (use 0.4 mL HCl from 12.1 N HCl; Fisher Scientific USA, to make up final volume 500 mL with distilled  $H_2O$ ).

- 2. Transfer buffer: 0.4 N NaOH, 1 M NaCl (weigh 16 g NaOH and 58.4 g NaCl and dissolve in distilled H<sub>2</sub>O to make up the final volume to 1000 mL).
  - 3. 20X SSC: 3M NaCl, 0.3 M sodium citrate trisodium salt (weigh 175.3 g NaCl, 88.2 g Na $_3$ C<sub>6</sub>H $_5$ O<sub>7</sub>.2H $_2$ O 900 mL H $_2$ O and adjust pH 7.0 using 1 N HCl and make up the final volume to 1000 mL with distilled H $_2$ O and autoclave).
    - 4. 2X SSC: Add 20 mL of 20X SSC in 180 mL of distilled  $H_2O$ . Protein analysis by Western blots.
    - 1. Acrylamide/Bis: ready made from Fischer (USA), stored at 4°C.
  - 2. 10% SDS: dissolve 10 g SDS in 90 mL deionized water, make up the volume to 100 mL, store at room temperature.
  - 3. Resolving gel buffer: 1.5 M Tris-HCl (add 27.23 g Tris base in 80 mL water, adjust to pH 8.8 with 6 N HCl and make up the final volume to 150 mL. Store at 4°C after autoclaving).
  - 4. Stacking gel buffer: 0.5 M Tris-HCl (add 6.0 g Tris base in 60 mL water. Adjust to pH 6.8 with 6 N HCl. Make up the volume to 100 mL. Store at 4°C after autoclaving).
- 5. Sample Buffer (SDS Reducing Buffer): In 3.55 mL water add 1.25 mL 0.5 M Tris-HCl (pH 6.8), 2.5 mL glycerol, 2.0 mL (10% SDS), 0.2 mL (0.5% Bromophenol blue). Store at room temperature. Add 50  $\mu$ L  $\beta$ -Mercaptoethanol ( $\beta$ ME) to 950  $\mu$ L sample buffer prior to its use.
- 6. 10X running buffer (pH 8.3): Dissolve 30.3 g Tris Base, 144.0 g
   25 Glycine and 10.0 g SDS in ~ 700 mL water (add more water if not dissolving). Bring up the volume to 1 L and store at 4°C.
  - 7. 10x PBS: Weigh 80 g NaCl, 2 g KCl, 26.8 g Na<sub>2</sub>HPO<sub>4</sub>·7 H<sub>2</sub>O (or 14.4 g Na<sub>2</sub>HPO<sub>4</sub>), 2.4 g KH<sub>2</sub>PO<sub>4</sub> in 800 mL water. Adjust pH to 7.4 with HCl and make up the volume to 1 L. Store at room temperature after autoclaving.
- 8. 20% APS: Dissolve 200 mg ammonium persulfate in 1 mL water (make fresh every two weeks).

9. Transfer buffer for 1500 mL: Add 300 mL 10x running buffer, 300 mL methanol, 0.15 g SDS in 900 mL water and make volume to 1 L.

## Plant Extraction Buffer:

		Used Concentration	Final Concentration
5	60 μ1	5M NaCl	100 mM
	60 μ1	0.5 M EDTA	10 mM
	600 μ1	1 M Tris-HCl	200 mM
	2 μ1	Tween-20	.05%
	30 μL	10% SDS	0.1%
10	$3~\mu L$	BME	14 mM
	1.2 mL	1 M Sucrose	400 mM
	1 mL	Water -	
	60 μL	100 mM PMSF	2mM

Add PMSF just before use (vortex to dissolve PMSF crystals).

PMSF (Phenylmethyl sulfonyl fluoride): Dissolve 17.4 mg of powdered PMSF in 1 mL of methanol by vortexing and store at -20°C for up to a month.

## Methods

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## Isolation of genomic DNA from plants.

Extract the genomic DNA from fresh green leaves using DNeasy Plant kit 20 (QIAGEN Inc.) following vender's instructions.

## Amplification of chloroplast flanking sequence.

Species-specific flanking sequences from the chloroplast DNA or genomic DNA of a particular plant species is amplified with the help of PCR using a set of primers that are designed using known and highly conserved sequence of the tobacco chloroplast genome.

Conditions for running PCR reaction: There are three major steps in a PCR, which are repeated for 30 to 40 cycles. (1) Denaturation at 94°C: to separate double stranded chloroplast DNA. (2) Annealing at 54 to 64°C: primers bind to single stranded DNA with formation of hydrogen bonds and the DNA polymerase starts copying the template. (3) Extension at 72°C: DNA Polymerase at 72°C extends to the template that strongly forms hydrogen bond with primers. Mismatched primers will not form strong hydrogen bonds and therefore, all these temperatures may vary based on

DNA sequence homology. The bases complementary to the template are coupled to the primer on the 3' side. The polymerase adds dNTPs from 5' to 3', reading the template in 3' to 5' direction and bases are added complementary to the template.

## Chloroplast transformation vector.

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The left and right flanks are the regions in the chloroplast genome that serve as homologous recombination sites for stable integration of transgenes. A strong promoter and the 5' UTR and 3' UTR are necessary for efficient transcription and translation of the transgenes within chloroplasts. For multiple gene expression, a single promoter may regulate the transcription of the operon, and individual ribosome binding sites must be engineered upstream of each coding sequence (2) (Fig. 10). The following steps are used in vector construction:

- 1. Amplification of flanking sequences of plastid with primers that are designed on the basis of known sequence of the tobacco chloroplast genome (between 16S-23S region of chloroplast).
- 2. Insert the PCR product containing the flanking sequence of the chloroplast genome into pUC19 plasmid digested with *PvuII* restriction enzyme (to eliminate the multiple cloning site), dephoshorylated with the help of alkaline phoshatase (CIP) for 5 min at 50°C (to prevent recircularization of cloning vector). Inactivate CIP enzyme at 68°C for 10 min.

Clone chloroplast transformation cassette (which is made blunt with the help of T4 DNA polymerase or Klenow filling) into a cloning vector digested at the unique PvuII site in the spacer region, which is conserved in all higher plants examined so far.

## Delivery of foreign genes into chloroplasts via particle gun.

This is most successful and a simple technique to deliver transgenes into plastids and is referred as Biolistic PDS-1000/ He Particle Delivery System (18,19). This technique has proven to be successful for delivery of foreign DNA to target tissues in a wide variety of plant species and integration of transgenes has been achieved in chloroplast genomes of tobacco (2), Arabidopsis (20), potato (21), tomato (25) and transient expression in wheat (22), carrot, marigold and red pepper (23) (see Note 5).

Preparation of gold particle suspension.

1. Suspend 50-60 mg gold particles in 1 mL 100% ethanol and vortex for 2 min.

- 2. Spin at maximum speed ~10, 000 x g (using tabletop microcentrifuge)
- 5 for 3 min.
  - 3. Discard the supernatant.
  - 4. Add 1ml fresh 70% ethanol and vortex for 1 min.
  - 5. Incubate at room temperature for 15 min and shake intermittently.
  - 6. Spin at 10, 000 x g for 2 min.
- 7. Discard supernatant, add 1ml sterile distilled H<sub>2</sub>O, vortex for 1min, leave at room temperature for 1min, and spin at 10, 000 x g for 2 min.
  - 8. Repeat above washing process three times with  $H_2O$  (step 7).
  - 9. Resuspend the gold-pellet in 1 mL 50% glycerol, store stock in -20°C freezer.

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Precipitation of the chloroplast vector on gold particles for five samples.

- 1. Take 50 µl the gold particles in 1.5 mL tube after vortexing for a 1 min.
- 2. Add 10  $\mu$ l DNA (about 1  $\mu$ g/ $\mu$ l plasmid DNA), and vortex the mixture for 30 sec.
- 3. Add 50 μl of 2.5 M CaCl<sub>2</sub> and vortex the mixture for 30 sec.
  - 4. Add 20  $\mu$ l of 0.1 M spermidine and vortex the mixture for 20 min at 4°C.

Washing of chloroplast vector coated on gold particles.

- 1. Add 200 μl 100% ethanol and vortex for 30 sec.
- 25 2. Spin at 3000 x g for 40 sec.
  - 3. Pour off ethanol supernatant.
  - 4. Repeat ethanol washings five times.
  - 5. In the last step, pour off ethanol carefully and add 35-40  $\mu$ l ethanol (100%).

Preparation of macrocarriers.

1. Sterilize macrocarriers by dipping in 100% ethanol for 15 min and insert them into sterile steel ring holder with the help of a plastic cap when airdried.

2. Vortex the gold-plasmid DNA suspension and pipet 8-10  $\mu$ l in the center of macrocarrier and let it air dry.

Gene gun setup for bombardment of samples.

- 1. Wipe the gun chamber and holders with 100% ethanol using fine tissue paper (do not wipe the door with alcohol).
  - 2. Turn on the vacuum pump.
- 3. Turn on the valve (Helium pressure regulator) of Helium gas tank (anti-clockwise).
- 4. Adjust the gauge valve (adjustable valve) ~200 to 250 psi above the desired rupture disk pressure (clockwise) using adjustment handle.
- 5. Turn on the gene gun.

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- 6. Place the rupture disc (sterilized by dipping in 100% ethanol for 5 min) in the rupture disc-retaining cap and tightly screw to the gas acceleration tube.
- 7. Place a stopping screen in the macrocarrier launch assembly and above that place macrocarrier with gold particles with chloroplast vector facing down towards screen. Screw assembly with a macrocarrier coverlid and insert in the gun chamber.
- 8. Place an intact leaf or explants to be bombarded on a filter paper (Whatman No. 1) soaked in medium containing no antibiotics. Place sample plate over target plate shelf, insert in the gun chamber and close the bombardment chamber door.
- 9. Press Vac switch to build pressure (up to 28 inches of Hg) in the vacuum gauge display. Turn same switch down at hold point and press Fire switch until you hear a burst sound of the ruptured disc.
- 10. Press Vent switch to release the vacuum and open the chamber to remove sample.
- 11. Shut down the system by closing the main valve (Helium pressure regulator) on the Helium gas cylinder. Create some vacuum in the gene gun chamber and keep using fire switch on and off until both pressure gauges' show zero reading. Release the vacuum pressure and turn off the gene gun and vacuum pump.

12. Incubate bombarded sample plates in the culture room for two days in the dark (i.e. covered with aluminum foil) and on the third day cut explants in appropriate pieces and place on the selection medium.

## Plant tissue culture and chloroplast transformation.

Tobacco chloroplast transformation.

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A highly efficient and reproducible protocol has been established for *Nicotiana tabacum* cv. Petit Havana (Daniell, H. (1997) *Methods in Mol. Biol. Recombinant gene expression protocols*. **62**,463-489.

- 1. Bombard 4 weeks old dark green tobacco leaves on the abaxial (bottom side) side with the chloroplast vector and incubate leaves in the dark for 2 days on selection free medium.
- 2. On the third day cut bombarded leaf explants into small square pieces (5 mm) and place explants facing abaxial surface towards selection medium containing MS salts, 1mg/l thiamine HCl, 100mg/l myo-inositol, 3% sucrose, 1mg/l BAP and 0.1 mg/l IAA along with 500 mg/l spectinomycin as a selective agent.
- 3. Transgenic shoots should appear after three to five weeks of transformation. Cut the shoot leaves again into small square explants (2 mm) and subject to a second round of selection for achieving homoplasmy on fresh medium.
- 4. Regenerate transgenic shoots (confirmed by PCR for transgene integration) on rooting medium containing MS salts, 1mg/l thiamine HCl, 100mg/l myo-inositol, 2% sucrose and 1mg/l IBA with 500mg/l spectinomycin.
- 5. Transfer transgenic plants into pots under high humidity and move them to green house or growth chamber for further growth and characterization.

Plastid transformation of edible crops.

The concept of universal vector for using the chloroplast DNA from one plant species to transform another species (of unknown sequence) was developed by the Daniell group (8). Using this concept both tomato and potato chloroplast genomes were transformed as described below.

Potato chloroplast transformation.

Using the tobacco chloroplast vector, leaf tissues of potato cultivar FL1607 was transformed via biolistics, and stable transgenic plants were recovered using the

selective aadA gene marker and the visual green fluorescent protein (GFP) reporter gene (21).

- 1. Bombard potato leaves (3-4 week old) and incubate in the dark for 2 days on selection free medium.
- 2. Third day excise leaves into small square pieces (5 mm) and place on MS medium containing B5 vitamins, 5 mg/l ZR, 0.1 NAA, and 3% sucrose. Gradually increase spectinomycin selection pressure (40 to 400 mg/l) after every two weeks subculture under diffuse light.
- 3. Regenerate shoots from transgenic potato calli on MS medium containing B5 vitamins, 0.01mg/L NAA, 0.1mg/L GA3, 2% sucrose and 40-400 mg/l spectinomycin.
  - 4. Transfer transgenic shoots on basal MS medium containing B5 vitamins, 2% sucrose and 40-400 mg/l spectinomycin for root induction. Transfer transgenic plantlets to growth chamber.

Tomato chloroplast transformation.

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Using the tobacco chloroplast vector, tomato (*Lycopersicon esculentum* cv. IAC Santa Clara) plants with transgenic plastids were generated using very low intensity of light (25).

- 1. Bombard four-week-old tomato leaves and incubate in the dark for 2 days on selection free medium.
  - 2. Excise bombarded leaves into small pieces and place on shoot induction medium containing 0.2 mg/L IAA, 3 mg/L BAP, 3% sucrose and 300 mg/L spectinomycin.
- 3. Select spectinomycin-resistant primary calli after a three to four month duration without any shoot induction.
  - 4. Regenerate shoots in about four weeks after transfer of transgenic calli to shoot induction medium containing 0.2 mg/L IAA, 2 mg/L ZR, 2% sucrose and 300 mg/L spectinomycin then root on hormone-free medium. Transfer regenerated transgenic plants into the greenhouse.

## 30 Molecular analysis of transgenic plants.

PCR screening of transgenic shoots.

This method has been used to distinguish between mutants, nuclear and chloroplast transgenic plants. By landing one primer on the native chloroplast genome adjacent to the point of integration and a second primer on the aadA gene (26. PCR product of an appropriate size should be generated in chloroplast transformants. Since this PCR product cannot be obtained in nuclear transgenic plants or mutants, the possibility of nuclear integration or mutants should be eliminated.

- 1) Extract the genomic DNA from transgenic leaf tissue using DNeasy Plant kit (QIAGEN Inc.) by following vender's instructions. For lower amount of transgenic tissues, volume of buffers may be reduced appropriately.
- 2) Run PCR reaction with Taq DNA Polymerase (QIAGEN Inc.) using appropriate primers following the same conditions as described above for amplification of flanking sequences.

Analysis of homoplasmy by Southern blot.

In Southern blot analysis, tobacco plastid genome digested with suitable restriction enzymes should produce a smaller fragment (flanking region only) in wild type plants compared to transgenic chloroplast that include transgene cassette as well as the flanking region. In addition, homoplasmy in transgenic plants is achieved when only the transgenic fragment is observed.

Transfer of DNA to membrane.

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- 1. Digest the genomic DNA (~2 to 10  $\mu$ g) with suitable restriction enzymes from transgenic samples (including wild type as a control) and run digested DNA on 0.8% agarose gel containing 5  $\mu$ L EtBr (from 10 mg/mL stock) in 100 mL for four hours at 40V.
- 2. Soak gel in 0.25 N HCl (depurination solution) for 15 minutes and rinse gel twice in distilled H<sub>2</sub>O for 5 minutes.
  - 3. Soak gel for 20 minutes in transfer buffer to denature DNA.
  - 4. Transfer overnight DNA from gel to nylon membrane (pre-soak first in water, then in transfer buffer for 5 minutes) using the transfer buffer.
- 5. Next day, rinse membrane twice with 2x SSC buffer for 5 minutes each and air-dry for 5 minutes on filter papers. Cross-link transferred DNA to membrane using GS GeneLinker UV Chamber (Bio-Rad) at appropriate (C3) setting.

Preparation of probe.

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1. Digest any plasmid (containing flanking sequences of the chloroplast genome) with appropriate restriction enzymes.

- 2. Denature 45  $\mu$ L flanking DNA fragment (50-250 ng) at 95°C for 5 minutes, then place on ice for 2-3 minutes.
  - 3. Add denatured probe to Ready-To-Go DNA Labeling Beads (-dCTP) tube (Amersham Biosciences, USA) and gently mix by flicking the tube.
  - 4. Add 5  $\mu$ L radioactive  $\alpha^{32}$ P (dCTP; Amersham Biosciences, USA) to probe mixture and incubate at 37°C for 1hour and filter the probe using ProbeQuant G-50 Micro Columns (Amersham Pharmacia Biotech Inc. USA).

Prehybridization and hybridization.

Place the blot (DNA transfer side facing towards the solution) in a hybridization bottle and add 10 mL Quik-Hyb (Stratagene, USA).

Incubate for 1 hour at 68°C. Add 100  $\mu$ L sonicated salmon sperm (10 mg/mL stock; Stratagene, USA) to the labeled probe and heat at 94°C for 5 minutes and add to bottle containing membrane and Quik-Hyb solution. Incubate for 1 hour at 68°C.

Washing and autoradiography.

- 1. Discard Quik-Hyb solution with probe and wash membrane twice in 50 mL (2x SSC buffer and 0.1% SDS) for 15 minutes at room temperature.
- 2. Wash membrane twice in 50 mL (0.1x SSC buffer and 0.1% SDS) for 15 minutes at 60°C.
- 3. Wrap the wash membrane in saran wrap and expose blot to x-ray film in the dark and leave at -70°C until ready for development.

Determination of transgene expression by Western blot.

Extraction of plant protein.

- 1. Grind 100 mg of leaf in liquid nitrogen and add 200  $\mu L$  of extraction buffer to samples on ice.
- 2. Add appropriate volume of freshly prepared 2x Sample loading buffer to an aliquot plant extract (from a stock containing 50  $\mu$ L  $\beta$ -mercaptoethanol and 950  $\mu$ L sample loading buffer).
- 3. Boil samples for 4 minutes with loading dye and centrifuge for 2 minutes at 10, 000 x g, then immediately load 20  $\mu$ L samples into gel.

Running gel.

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Load samples on gel and run for half hour at 100 V, then 1 hour at 150 V until the marker bands corresponding to your protein are in middle.

Transfer of protein to membrane.

Transfer protein from gel to membrane using Mini Transfer Blot Module at 30 V overnight or 65 V for 2 hours or 100 V for 1 hour. Membrane wrapped in saran wrap can be stored at -20°C for a few days if necessary.

## Membrane blocking

- 1. After transfer, rinse membrane with water and incubate membrane in PTM (100 mL 1x PBS, 50  $\mu$ L 0.05% Tween 20, and 3 g dry milk (3%) for 1 hour at room temperature.
  - 2. Add primary antibody in suitable dilution for 15 mL and incubate for 2 hours at room temperature. Wash membrane twice with 1x PBS for 5 minutes each.
  - 3. Add secondary antibody in proper dilution for 20 mL. Incubate for 1.5 hours at room temperature on a shaker.
  - 4. Wash twice with PT (100 ml 1x PBS + 50  $\mu$ L Tween 20) for 15 minutes and finally with 1x PBS for 10 minutes.

## Exposure of the blot to X-ray film.

- 1. Mix 750 μL of each chemiluminescent solution (Luminol Enhancer and Stable Peroxide) in 1.5 mL tube and add to membrane, cover thoroughly.
  - 2. Wipe out extra solution and expose blot to x-ray film for appropriate duration and develop film.

#### Seed sterilization.

- 25 1. Vortex small amount of seeds into microcentrifuge tube with 1 mL 70% ethanol for 1 minute. Discard ethanol after brief spin.
  - 2. Add 1 mL disinfecting solution (1.5% Bleach and 0.1% Tween 20) in tube and vortex intermittently for 15 min. Discard solution after brief spin.
    - 3. Wash the seed thrice with sterile distilled water.
- 4. Spray seeds with sterile water on plate containing RMOP basal medium supplemented with 500  $\mu$ g/mL spectinomycin to determine maternal inheritance in transgenic chloroplast plants.

## Evaluation of results.

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Maternal inheritance in chloroplast transgenic plants.

Transgenes integrated into chloroplast genomes are inherited maternally. This is evident when transgenic seed of tobacco are germinated on RMOP basal medium containing 500  $\mu$ g/mL spectinomycin. There should be no detrimental effect of the selection agent in transgenic seedlings whereas untransformed seedlings will be affected.

CTB-GM1-gangliosides binding ELISA assay.

- Coat microtiter plate (96 well ELISA plate) with monosialoganglioside GM1 {3.0 μg/mL in bicarbonate buffer (15 mM Na<sub>2</sub>CO<sub>3</sub>, 35 mM NaHCO<sub>3</sub>, pH 9.6)} and as a control, coat BSA (3.0 ug/mL in bicarbonate buffer) in few wells.
  - 2. Incubate plate overnight at 4°C.
  - 3. Block wells with 1% (w/v) bovine serum albumin (BSA) in 0.01 M phosphate-buffered saline (PBS) for two hours at 37°C.
    - 4. Wash wells thrice with PBST buffer (PBS containing 0.05% Tween 20).
  - 5. Incubate plate by adding soluble protein from transformed and untransformed plants and bacterial CTB in PBS.
  - 6. Add primary antibodies (rabbit anti cholera serum diluted 1:8000 in 0.01 M PBST containing 0.5% BSA) and incubate plate for 2 hours at 37°C.
    - 7. Wash well thrice with PBST buffer.
  - 8. Add secondary antibodies diluted 1:50,000 (mouse anti rabbit IgGalkaline phosphatase conjugate in 0.01 M PBST containing 0.5% BSA) and incubate plate for 2 hours at 37°C.
- 9. Develop plate with Sigma Fast pNPP substrate. Stop reaction by adding 3 M NaOH and read plate absorbance at 405 nm.

The macrophage lysis assay is as follows:

- 1. Isolate crude extract protein from 100 mg transgenic leaf using 200  $\mu$ L of extraction buffer containing CHAPS detergent (4% CHAPS, 10 mM EDTA, 100 mM NaCl, 200 mM Tris-HCI, pH 8.0, 400 mM sucrose, 14 mM  $\beta$ -mercaptoethanol, 2 mM PMSF) and one without CHAPS detergent.
- 2. Spin samples for five minutes at 10, 000 x g and use both supernatant and homogenate for assay

3. Plate macrophage cells RAW 264.7 (grown to 50% confluence) into 96-wells plate, incubated in 120  $\mu$ L Dulbecco's Modified Eagle's Medium (DMEM; from Invitrogen life technologies).

- 4. Aspirate medium from wells and add 100 μL medium containing 250
   5 ng/mL proteins in crude leaf extract.
  - 5. In control plate, add only DMEM with no leaf fraction to test toxicity of plant material and buffers.
  - 6. In another plate, add 40  $\mu$ L dilutions onto RAW 264.7 cells from plant samples, which serially diluted 2 fold, so that the top row had plant extract at 1:14 dilution.
  - 7. Add 20 □L of MTT 3-[4,5-dimethylthiazol-2-yl]-2,5-diphenyltetrazolium bromide; Sigma) to each well containing cells (from a stock 5mg/ml MTT dissolved in 1xPBS and filter sterilize) after 5 hours to assess the cell death.
  - 8. Incubate the plate at 37°C for 5 hours. Remove media with needle and syringe. Add 200 □L of DMSO to each well and pipette up and down to dissolve crystals. Transfer to plate reader and measure absorbance at 550nm.

Active PA was found in both the supernatant and homogenate fractions. However, maximum macrophage lysis activity was noticed in supernatant when extraction buffer was used with CHAPS detergent.

Cholera toxin (CTB) antigen as an edible vaccine.

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Chloroplast transgenic plants are ideal for production of vaccines. The heatlabile toxin B subunits of E. coli enterotoxin (LTB), or cholera toxin of Vibrio cholerae (CTB) have been considered as potential candidates for vaccine antigens. Integration of the unmodified native CTB gene into the chloroplast genome has demonstrated high levels of CTB accumulation in transgenic chloroplasts (Daniell, H., et al. (2001). J. Mol. Biol. 311,1001-1009.). This new approach not only allowed the high level expression of native CTB gene but also enabled the multimeric proteins to be assembled properly in the chloroplast, which is essential because of the critical role of quaternary structure for the function of many vaccine antigens. The expression level of CTB in transgenic plants was between 3.5% and 4.1% tsp and the functionality of the protein was demonstrated by binding aggregates of assembled pentamers in plant

extracts similar to purified bacterial antigen, and binding assays confirmed that both chloroplast-synthesized and bacterial CTB bind to the intestinal membrane GM1-ganglioside receptor, confirming correct folding and disulfide bond formation of CTB pentamers within transgenic chloroplasts (Fig. 11).

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Further, this invention contemplates the examples of edible vaccines expressed via the plastid, such as CTB( as described above), Anthrax, Plague, and all other vaccines known and desribed in the art including those described in PCT/US02/41503, filed on 12/26/02. The aforementioned application is fully incorporated by reference. The aspects of this invention further contemplate the expression of other therapeutic proteins such as interferon, IGF-1, insulin, which will be expressed in non-green plant cells for oral delivery. An exemplary description of such theraputic proteins can be found in the PCT applications which claim priority to U.S. provisional application Nos. 60/393,439;60/393,428;60/393,451; 60/393,649; and 60/393,438, wherein the expression of theraputic proteins in the plant plastid is described.

Oral delivery of vaccines and selection of transgenic plants without the use of antibiotic selectable markers.

Betaine aldehyde dehydrogenase (*BADH*) gene from spinach has been used as a selectable marker to transform the chloroplast genome of tobacco (Daniell, H. et al., (2001) *Curr. Genet.* **39**,109-116). Transgenic plants were selected on media containing betaine aldehyde (BA). Transgenic chloroplasts carrying BADH activity convert toxic BA to the beneficial glycine betaine (GB). Tobacco leaves bombarded with a construct containing both *aadA* and *BADH* genes showed very dramatic differences in the efficiency of shoot regeneration. Transformation and regeneration was 25% more efficient with BA selection, and plant propagation was more rapid on BA in comparison to spectinomycin. Chloroplast transgenic plants showed 15 to 18 fold higher BADH activity at different developmental stages than untransformed controls. Expression of high BADH level and resultant accumulation of glycine betaine did not result in any pleiotropic effects and transgenic plants were morphologically normal and set seeds as untransformed control plants.

# Production of human therapeutic proteins in transgenic chloroplasts.

Human serum albumin (HSA) protein.

Human Serum Albumin (HSA) accounts for 60% of the total protein in blood and widely used in a number of human therapies. Chloroplast transgenic plants were generated expressing HSA (Fernandez-San Millan et al., (2003) *Plant Bitechnol. J.* 1,71-79). Levels of HSA expression in chloroplast transgenic plants was achieved up to 11.1% tsp. Formation of HSA inclusion bodies within transgenic chloroplasts was advantageous for purification of protein. Inclusion bodies were precipitated by centrifugation and separated easily from the majority of cellular proteins present in the soluble fraction with a single centrifugation step. Purification of inclusion bodies by centrifugation may eliminate the need for expensive affinity columns or chromatographic techniques.

# Purification of HSA.

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- 1. Solubilize the HSA inclusion bodies from transformed tissues using extraction buffer containing 0.2M NaCl, 25 mM Tris-HCl (pH 7.4), 2mM PMSF and 0.1% Triton X-100.
- 2. Spin at 10, 000 x g. Suspend the pellet in buffer containing 6M Gu-HCl, 0.1M  $\beta$ ME and 0.25 mM Tris-HCl (pH 7.4).
- 3. Dilute plant extract 100-fold in buffer containing 100 mM NaCl, 50 mM Tris-HCl (pH 8.5) and 1 mM EDTA.
  - 4. Concentrate HSA protein by precipitation using a polyethylenglycol treatment at 37%.
  - 5. Separate protein fractions by running a SDS-PAGE gel and stain gel with silver regent following vender's instruction (Bio-Rad, USA).

Electron microscopy and immunogold labeling.

- 1. Cut the transformed and untransformed leaf in 1-3 mm squares.
- 2. Fix them in 0.1 M cacodylate buffer pH 7.4 (2.5% glutaraldehyde, 2% paraformaldehyde and 5 mM CaCl<sub>2</sub>) for 15 minutes under vacuum and 12 hours at 4°C.
  - 3. Rinse samples twice in 0.1M cacodylate buffer (pH 7.4) after fixation.
- 4. Dehydrate fixed samples through a graded ethanol series to 95%, then implant in LRW resin at 60°C for 24 hours.

5. Cut ultra-thin sections using a Leica Ultracut T ultramicrotome and collect sections onto nickel grids.

- 6. Incubate sections in 0.05M glycine prepared in PBS buffer for 15 minutes to inactivate residual aldehyde groups.
- 7. Place grids onto drops of blocking solution (PBS containing 2% non-fat dry milk) and incubate for 30 minutes
- 8. Incubate sections for 1 hour in a goat anti-human albumin polyclonal antibody (dilution range from 1:1000 to 1:10,000 in blocking solution).
  - 9. Wash sections with blocking solution 6 X 5 minutes each.
- 10. Incubate sections for 2 hours with a rabbit anti-goat IgG secondary antibody conjugate to 10 nm gold diluted 1:40 in blocking solution.
- 11. Wash sections 6 X 5 minutes in blocking solution and 3 X 5 minutes with PBS, and fixed sections in 2% glutaraldehyde diluted in PBS for 5 minutes.
- 12. Wash fixed sections in PBS 3 X 5 minutes, then in distilled water 5 X 2 min each.
- 13. Stain sections using uranyl acetate and lead citrate and examine samples under transmission electron microscope at 60kv.

## Notes

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- 1. Gold particles suspended in 50% glycerol may be stored for several months at -20°C. Avoid refreezing and thawing spermidine stock; use once after thawing and discard the remaining solution. Use freshly prepared CaCl<sub>2</sub> solution after filter sterilization. Do not autoclave.
  - 2. Precipitation efficiency of DNA on gold and spreading of DNA-gold particles mixture on macrocarriers is very important. For high transformation efficiency via biolistics, a thick film of gold particles should appear on macrocarrier disks after alcohol evaporation. Scattered or poor gold precipitation reduces the transformation efficiency.
  - 3. Generally, a 1000 bp flanking sequence region on each side of the expression cassette is adequate to facilitate stable integration of transgenes.
  - 4. Use of the 5' untranslated region (5' UTR) and the 3' untranslated region (3' UTR) regulatory signals are necessary for higher levels of transgene expression in plastids (13). The expression of transgene in the plant chloroplast depends on a

functional promoter, stable mRNA, efficient ribosomal binding sites; efficient translation is determined by the 5' and 3' untranslated regions (UTR). Chloroplast transformation elements *Prrn*, *psbA5'UTR*, *3'UTR* can be amplified from tobacco chloroplast genome.

5. Bombarded leaves after two-days dark incubation should be excised in small square pieces (5-7 mm) for first round of selection and regenerated transgenic shoots should be excised into small square pieces (2-4 mm) for a second round of selection.

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- 6. Temperature for plant growth chamber should be around 26-28°C for appropriate growth of tobacco, potato and tomato tissue culture. Initial transgenic shoot induction in potato and tomato require diffuse light. However, higher intensity is not harmful for tobacco.
  - 7. Transformation efficiency is very poor for both potato and tomato cultivars compared to tobacco.
  - 8. Tobacco chloroplast vector gives low frequency of transformation if used for other plant species. For example, when petunia chloroplast flanking sequences were used to transform the tobacco chloroplast genome (DeGray, G. et al., (2001), *Plant Physiol.* 127,852-862.), it resulted in very low transformation efficiency.

Under diffuse light conditions, highly regenerating tomato cultivar (Microtom) shoots produce premature flowering that inhibit further growth of transgenic plants. Therefore, after the first shoot induction phase, shoots should be moved to normal light conditions.

# ILLUSTRATIVE EXAMPLE 1: SOMATIC EMBRYOGENESIS VIA CARROT TRANSFORMATION

Homoplasmic transgenic carrot plants exhibiting high levels of salt tolerance (up to 500 mM NaCl) were rapidly regenerated from carrot cell cultures, via somatic embryogenesis. Carrot chloroplast genome is strictly maternally inherited and plants do not produce seeds in the first year, offering complete containment of transgene flow. Carrot cells multiply rapidly and large biomass is produced using bioreactors; somatic embryos are derived from single cells; viable for long duration on culture medium, encapsulated embryos are used as synthetic seeds for cryopreservation and controlled

germination; these features provide an ideal production system for plant made pharmaceutical proteins and their oral delivery. BADH expressing cells offer a visual selection by their green color, distinguishing them from untransformed yellow cells. A useful trait has been engineered via the chloroplast genome for the first time in a non-tobacco crop. This is the first plastid transformation achieved using non-green explants and somatic embryogenesis, opening the door to transform monocots, legumes, vegetables, fruit crops and transgene expression in non-green edible parts.

This Example presents the first report of stable and highly efficient plastid transformation of carrot using non-green tissues as explants, regenerated via somatic embryogenesis. An useful plant trait (salt tolerance) has been expressed for the first time in a non-solanaceous crop via the chloroplast genome. So far, only the tobacco chloroplast genome has been engineered to confer herbicide resistance (Daniell et al., 1998), insect resistance (McBride et al., 1995; Kota et al., 1999; DeCosa et al., 2001), disease resistance (DeGray et al., 2001), drought tolerance (Lee et al., 2003) and phytoremediation of toxic metals (Ruiz et al., 2003). In addition, expression of the BADH enzyme facilitated the visual selection of transgenic green cells from non-transgenic yellow cells. As a major crop consumed world-wide, this report should serve as a model for genetic engineering of plastid genomes in higher plant species that require the use of non-green tissues as explants and somatic embryogenesis for regeneration.

# **Results and Discussion**

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## Construction of carrot plastid transformation vectors.

Carrot chloroplast vectors target the expression cassette to the 16S/trnI – trnA/23S region of the chloroplast genome for integration via homologous recombination. The site of integration is similar to the universal chloroplast transformation vector (pLD) reported earlier from our laboratory (Daniell et al., 1998; Guda et al., 2000) but the size of the flanking sequence on either side of the expression cassette was doubled to enhance efficiency of homologous recombination. As a result the flanking sequence were increased to approximately 4kb. The chloroplast transformation vector pDD-Dc-gfp/BADH (Fig. 1A) is a carrot specific vector that harbors the gfp gene regulated by the gene 10 5'UTR / rps16 3'UTR (to facilitate expression in green as well as non-green tissues and screen transformants by GFP

fluorescence) and the *badh* gene expressed under the *psbA* 5' and 3' UTRs (to facilitate expression in green tissues, regulated by light). All the 5' and 3' regulatory elements were PCR amplified from tobacco genomic DNA except for the gene 10 5'UTR which was derived from phage T7 gene 10. Transcription of both transgenes in this cassette is driven by the full length *Prrn* 16S rRNA promoter which contains regulatory elements for both the nuclear encoded and plastid encoded RNA polymerases, thereby facilitating transcription in green as well as non-green tissues. The chloroplast transformation vector pDD-*Dc-aadA/BADH* (Fig. 1B) harbors the *aadA* and *badh* genes whose expression is driven by the full length 16S rRNA promoter under the regulation of RBS (Ribosome Binding Site) / 3'psbA UTR and the gene 10 5'UTR / rps16 3'UTR regulatory elements, respectively.

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It should be further noted that the universal vector described in U.S. patent Application No. 09/079,640 can be used to transform the plastid genome as described within this application.

# Transformation of carrot plastids and plant regeneration.

Yellow fine cell suspension culture induced from stem segments of carrot (Daucus carota L. cv. Half long) was bombarded with carrot chloroplast vectors pDD-Dc-gfp/BADH and pDD-Dc-aadA/BADH as described (Daniell, 1997; Kumar and Daniell, 2003). Carrot cultures transformed with pDD-Dc-gfp/BADH vector, produced GFP expressing calli and somatic embryos that were observed under confocal microscope (Fig. 2 A-C) when selected on 20 mM betaine aldehyde (BA). Even though it was quite evident that somatic embryos expressing GFP could be regenerated rapidly on BA selection, there were two major limitations for using this vector. BA is quite expensive (\$2000-\$3000/ gm) and this limited the number of experiments that could be performed and the badh gene was regulated by the psbA promoter and UTR elements, that was under developmental and light regulation. Unfortunately, cultured carrot cells were non-green and in early stages of development, thus minimizing badh expression. Therefore, more efforts were made to transform carrot cells using another chloroplast vector. Using the chloroplast vector pDD-Dc-aadA/BADH, seven independent transgenic cell lines were recovered within 2-3 months from five bombardments on solid medium (MSB+ 3 mg/L 2,4-D, 1 mg/L kinetin) containing different concentrations of spectinomycin (150-450 mg/L). Further, transgenic calli were

transferred to 350-mg/L spectinomycin for a month and multiplied using 500-mg/L spectinomycin. All Transgenic cell cultures were incubated under 100 lx light intensity at 26±2°C temperature. The MSB medium supplemented with 3-mg/L 2,4-D and 1 mg/L kinetin along with the selection agent was used as the growth medium to induce and multiply the transgenic cell cultures. In order to multiply transgenic cultures, they were subcultured on solid medium every 2-3 week and were grown in liquid medium (MSB+0.1 mg/L 2,4-D) at 130 rpm under diffuse light (50 lx). The medium without 2,4-D (MSB+0.2 mg/L kinetin) was used as the plant-regeneration-medium to convert embryogenic calli into plantlets. Transgenic plants maintained on basal MSB medium (containing 500 mg/L spectinomycin) were transferred to soil in pots to induce the mature taproot system and used for further characterization.

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Visible selection of transgenic carrot cells. During in vitro cell culture studies of transgenic and non-transgenic carrot, it was interesting to note that carrot cells could be distinguished on the basis of color. Transgenic carrot cells, which carry the badh transgene, were always green in color whereas non-transgenic cells were yellow in color (see Fig. 3A-B). To test that transgenic bright green cells were truly transgenic, hetroplasmic (partially transformed plastids) carrot cell cultures were placed on a growth medium without selection and were allowed to segregate; green and yellow cells visually segregated within 3-4 weeks (Fig. 3C-D). Transgenic-green cells were confirmed positive for transgene integration while yellow cells were found to be untransformed. Further, suppression of untransformed cell division was possible, when cells were exposed to different concentrations of NaCl. This visible selection system would be vital to distinguish transformed from untransformed cells while eliminating the selectable antibiotic marker genes using direct repeats (Iamtham and Day, 2002) from transformed chloroplast genomes. It will be interesting to investigate the role of BADH enzyme in enhancing chlorophyll biosynthesis or stability in transgenic chloroplasts.

# Confirmation of transgenes integration into carrot plastids.

The carrot chloroplast vector pDD-Dc-aadA/BADH integrates the aadA and badh genes into the 16S-23S-spacer region of the plastid genome by homologous recombination. Transgene integration into carrot cell lines was confirmed by PCR (Fig. 4A) using internal primer set 3P (that lands on trnI region of plastids) and 3M (that

lands on the *aadA* gene) producing 1.6 kb PCR product. This eliminates mutants that may be obtained on spectinomycin selection, caused by mutation of the chloroplast 16S rRNA gene. In order to distinguish between nuclear and chloroplast transgenic cell lines (Fig. 4B), 16S-F primer was landed on the native chloroplast genome, 200 bp upstream of integration site and 1M primer was landed on the *aadA* gene; this generated 2.5 kb size PCR product. Since this PCR product cannot be obtained in nuclear transgenic plants, the possibility of nuclear integration was eliminated. Thus, PCR analyses allow rapid screening of large number of putative transgenic lines and eliminate mutants or nuclear transgenic lines.

# Determination of homoplasmy in transgenic carrot plastids.

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PCR confirmed transgenic carrot cell lines were repeatedly subcultured each week in liquid medium for several rounds of selection in the presence of selection agent. Southern blot analysis was performed using total genomic DNA isolated from untransformed and transformed carrot plants, developed from different transgenic cell lines and was digested with AfIIII and PvuII (Fig. 4C). Presence of AfIIII restriction site in the 16S-rRNA region (left flank) in both transformed and untransformed carrot plastids and a unique PvuII site in between trnI and trnA flanking regions of untransformed plastids as well as in the mid region of badh transgene, allowed excision of predicted size fragments in both the untransformed and transgenic lines. In order to confirm heteroplasmy or homoplasmy, genomic DNA of carrot plants digested with AfIIII and PvuII was hybridized with the 4.9 kb radioactive DNA probe. This probe fragment was isolated from the chloroplast vector pDD-Dc-aadA/BADH, by digesting with AfIIII and PvuII; this fragment includes the 2.4 kb trnI flanking sequence and 2.5 kb transgene sequences of the chloroplast vector (Fig. 1B). Transgenic plants developed after two subcultures in liquid medium on the selection agent (350 mg/L spectinomycin) showed heteroplasmy (Fig. 4C, lane 2), whereas plants that were developed from cell lines after 8-10 subcultures in liquid medium supplemented with high concentration of selecting agent (500 mg/L spectinomycin) showed homoplasmy (Fig. 4C, lanes 3-8).

# BADH enzyme activity in carrot cells, root and shoot.

BADH enzyme activity was assayed in crude extracts from untransformed and transformed carrot cell cultures, tap roots (carrot) and leaves as described (Daniell et al

2001). By assessing BADH enzyme activity, the expression of badh transgene was observed in cells and different parts of carrot plants, using 50 µg crude extract of protein from each sample, desalted using G-25 column. BADH enzyme in the presence of betaine aldehyde converts NAD<sup>+</sup> to NADH and the reaction rate was measured by an increase in absorbance at 340 nm due to the reduction of NAD<sup>+</sup>. Crude extracts from transgenic plastids (cells, tap roots and leaves) demonstrate elevated levels of BADH activity compared to untransformed tissues of carrot (Fig. 5A). Higher BADH activity was observed in leaves, tap roots of carrot plant and transgenic cells in suspension culture, confirming that full length 16S promoter Prrn and gene 10 UTR are highly suitable for expressing transgenes in various tissues. Because these regulatory elements are anticipated to function uniformly in all tissues, we presume that the difference in BADH activity observed might be due to variation in the plastid genome copy numbers. It is known that plastid genome copy numbers vary significantly in different tissues, with only 5% observed in roots compared to leaves (Sasaki et al 1990). However, the high BADH enzyme activity observed in carrot tap root (75% of leaves) may be due to the large number of chromoplasts present; this was evident by their orange or purple color, instead of colorless roots normally observed in plants that contain only 5% of plastid copies.

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# BADH protein expression in carrot cells, root and shoot.

To further confirm the results of BADH activity in cells, tap roots and leaves, western blot analysis was done using crude extracts prepared from transformed and untransformed carrot tissues and a fraction from each sample (50 µg protein) was subjected to 10% SDS-PAGE. Protein transferred to nitrocellulose membrane was hybridized with polyclonal anti-BADH serum, raised in rabbits against native BADH (kindly provided by Dr. Elisa Soto; Figueroa-Soto et al 1999) and antigenic peptides were detected using horseradish peroxidase-linked secondary antibody. No *badh* expression was detected in untransformed carrot tissues (cells, tap roots and leaves; lanes 1-3). However, in transgenic samples (Fig. 5B), higher expression was observed in leaves (lane 6) and tap roots (lane 7) compared to carrot cell suspension cultures (lane 4). BADH protein accumulation in carrot root and leaf tissues were parallel to the results obtained with BADH enzyme activity in transgenic roots and shoots.

# Salt tolerance and BADH activity in cell suspension cultures of carrot.

In order to test whether salt stress affects the BADH activity, experiments were performed under different salt concentrations (0-500 mM NaCl) on carrot cell suspension cultures. Study performed for two weeks on carrot cells showed that transformed cells were able to sustain and proliferate in higher concentrations of NaCl in the liquid medium than untransformed cells (Fig. 6A-B). Using 500 mL conical flask containing 100 ml medium, carrot cultures produced about 11. 82 g cells in both transformed and untransformed (1475%) while, in presence of 100 mM NaCl, 8.75 g transgenic cells (1096%) and 1.29 g untransformed cells (161%) were obtained after two weeks (when initial culture used contained 0.8 g per flask, see Fig. 6A-B). Furthermore, BADH enzyme activity was stimulated in transgenic carrot cell cultures when they were exposed to 100 to 300 mM NaCl. Maximum noticeable BADH activity was seen in 100 and 200 mM NaCl (Fig. 6C) and such increase was insignificant in untransformed cells. This shows that full length Prrn promoter and gene 10 5' UTR facilitate efficient transcription and translation in all tissues, irrespective of the developmental stage, despite low copy number of plastid genomes in non-green cells or roots.

## Effect of salt on carrot plants.

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Different salt concentrations (100–500 mM NaCl) were tested on transformed and untransformed carrot plants transferred to soil in pots. Transgenic plants carrying the *badh* transgene thrive well up to 400 mM NaCl (Fig. 7) whereas, untransformed plants could not survive in pots after two weeks over 200 mM NaCl.

One aspect of this invention describes methods for transforming plastids using a highly efficient process for carrot plastid transformation through somatic embryogenesis. Still other aspects of this invention provide for vectors which are capable plastid transformation through somatic embryogenesis. Still another aspect provides for transformed plastids, plants, and plant parts, which have been transformed through somatic embryogenesis through the methods and vectors described herein. This application, along with the knowledge of the art, provides the necessary guidance and instructions to engineer the plastid genome of several major crops in which regeneration is mediated through somatic embryogenesis. Cereal crops (wheat, rice, corn, sugarcane), legumes (soybean, alfalfa), oil crops (sunflower, olive), cash crops

(cotton, coffee, tea, rubber, flex, cork oak, pines), vegetables (eggplant, cucumber, cassava, chili pepper, asparagus etc.), fruits (apple, cherry, banana, plantain, melons, grape, guava), nuts (cashew, walnuts, peanuts), and trees (date palm etc.) are regenerated through somatic embryogenesis.

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Previously, chloroplast transformation has been reported through bombardment of large number of leaf explants to obtain few transgenics. For example, in potato out of 200 bombardments only 2 transgenic shoots were recovered (Sidorov, 1999) after prolonged tissue culture. Authors reported that transformation of tomato plastids took almost two years (Ruf etal 2001). In contrast, stable transgene integration was confirmed in seven transgenic cell lines of carrot within 8-10 weeks from five bombardments and homolplasmy was achieved in cell cultures by repetitive 8-10 weekly subcultures in liquid medium. Carrot transgenic plants (ready to go in pots) were generated within a month from homoplasmic cell lines. Such high frequency of transformation obtained in carrot for transgene integration into plastid genomes might be due to several reasons. Long flanking sequences for homologous recombination were used from the same species i.e. native chloroplast genome of carrot (Daucus carota L. cv. Half long). In contrast, chloroplast vectors used for plastid transformation in both tomato and potato contained flanking sequences from the tobacco chloroplast genome. Similarly, when petunia chloroplast flanking sequences were used to transform the tobacco chloroplast genome (DeGray et al 2001), it resulted in very low transformation efficiency. Therefore, in order to advance this field of research, chloroplast genomes of useful crops should be sequenced. It is quite challenging to obtain flanking sequences for homologous recombination when no chloroplast DNA sequence is available in the Genbank (as it was the case for carrot).

Another significant observation in this study is high levels of transgene expression in proplastids of cultured carrot cells. Sidorov et al reported 50-fold less GFP accumulation in amyloplasts of potato tubers compared to leaves. In sharp contrast we report 53% BADH activity observe in cells comparison to leaves. This is the first report of transgene expression in proplastids even though chromoplasts have been reported to accumulate reasonable amounts of foreign gene products (Ruf et al 2001). The heterologous gene 10 5' UTR that regulates translation of *badh* gene was indeed promoterless. Chloroplast psbA 3' UTR is known to stabilize transcripts and is

poor in transcription termination, with 30% termination efficiency. However, if higher levels of expression are desired in proplastids (such as corn seeds or in roots for nematode resistance), it may be desirable to insert a promoter upstream of gene 10 5'UTR to enhance transcription.

In order to achieve homoplasmy in carrot system, several rounds of selection (8-10) were required along with high selection pressure. In liquid culture, all surfaces of carrot cells are directly exposed to the growth medium. This implies a high selection pressure, which might possibly block the growth of untransformed plastids. Carrot cells in liquid medium multiply fast compared to solid medium containing 2,4-D. Once 2,4-D is removed from the carrot cells, they rapidly convert into mature somatic embryos and profusely give rise to plantlets. Yet another requirement to achieve homoplasmy is the availability of templates for integration into thousands of plastid genomes per cell. We have previously shown that the use of chloroplast flanking sequence that contain the complete chloroplast origin of replication offers large number of templates within chloroplasts (Daniell et al., 1990) and helps to achieve homoplasmy even in the first round of selection (Guda et al., 2000). Inclusion of the chloroplast origin of replication within the carrot chloroplast vectors (assuming that the *ori* is present in the same location as in other plant species) should have helped achieve homoplasmy within a few rounds of cell division.

Carrot cells have a great potential for rapid proliferation through somatic embryos, which produce secondary somatic embryos (recurrent embryogenesis) until exposed to maturation medium for conversion into plants and this culture can be maintained for several year in vitro culture. In *Daucus carota* L. cv. *US-Harumakigosun*, embryogenic carrot calli maintained on suitable growth medium for five year produced a large number of plants (4×10<sup>7</sup> plantlets/*L*-medium/day) continuously for 245 days without any decrease in the productivity (Nagamori etal 1999) through liquid cell culture using a rotating bioreactor. Another advantage with somatic embryos is that they can be used as synthetic seeds. Synthetic or artificial seeds have been defined as somatic embryos encapsulated in sodium aliginate beads or matrix for use in the commercial propagation of plants. For example, carrot somatic embryos encapsulated in alginate-gellan gum (dehydrated to 15% RH and rehydrated in moistured air to 90% RH) germinated up to 73% in soil (Timbert etal 1996) and the

frequency of immobilized cell regeneration was improved about 1.5 times higher than that obtained through non-immobilized cells (Nagamori et al 1999). With synthetic seed technology, desirable elite genotypes in carrot and other plant species can be cryopreserved for long time (Tessereau et al 1994) and on demand these propagules can be used for synchronized planting of plants in the greenhouse or field. A single source for production in highly desired to minimize heterogeneity of therapeutic proteins.

Glycine betaine (GB) is a commonly occurring compound in all-living organisms and studies suggests that many higher plants, bacteria, marine and mammalian species accumulate GB under water deficiency or salt stress. In plants, bacteria and in mammalian kidney tissues, GB acts as an osmoprotectant. In porcine kidney BADH is localized in cortex and in the inner medulla, which is specifically localized on cells surrounding the tubules that are exposed to urea and high salt compounds (Figueroa-Soto et al 1999). Glycine betaine is one of the major compatible organic osmolytes identified in renal inner medulla of rats, which help renal cells to survive and function normally, despite being exposed to hypertonicity and lethal levels of urea.

These facts suggest that the presence of the BADH enzyme or accumulation of GB would be beneficial in foods consumed by humans or animals. In plants, salinity stress is a continuing and increasingly deleterious to the growth and yield of crop plants, owing to irrigation practices and increasing demands on fresh water supply; therefore, engineering of salt-tolerant crop plants has been a long-held and intensively sought objective (Apse MP and Blumwald E, 2002). One effective mechanism to reduce damage from these stresses is the accumulation of high intracellular levels of osmoprotectant compounds like glycine betaine in plants through genetic engineering (Rontein etal 2002).

## Experimental protocol.

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# Construction of plastid transformation vectors.

DNA fragment representing carrot flanking sequence was amplified from carrot genomic DNA that was isolated from the leaves using DNeasy Plant mini kit (Qiagen Inc.) following manufacturer's protocol. The flanking sequence fragment was amplified with the primers generated from the tobacco chloroplast genome sequence, using Platinum Pfx DNA polymerase (Invitrogen Inc.). The amplified fragment

represents the 16S/trnI-trnA/23S region of the chloroplast genome and is approximately 4.2 kb in size. The PCR amplified DNA fragment was treated with T4 polynucleotide kinase (Promega) and cloned into PvuII digested pBluescript II KS, dephosphorylated with Shrimp Alakaline phosophatase (Promega). The kinase and dephosphorylation reactions were performed as per the manufacturer's instructions. The chloroplast promoters and regulatory sequences were amplified using PCR based on the information available for the tobacco chloroplast genome (Accession number -NC 001879). The carrot specific chloroplast transformation vector pDD-Dcgfp/BADH (Fig. 1A) was constructed by inserting a blunt ended ClaI-SacI fragment representing the gfp/BADH expression cassette into PvuII site of carrot chloroplast DNA flanking sequences. The sm-gfp gene was obtained from TAIR. Similarly, the carrot chloroplast transformation vector pDD-Dc-aadA/BADH (Fig. 1B) was constructed by inserting blunt ended ApaI fragment representing the aadA/BADH expression cassette into carrot chloroplast DNA flanking sequences at the PvuII site, after dephoshorylation. Bacterial and DNA manipulations were performed as per standard molecular biology protocols.

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Of note, a potential vector for use is the double barrel plastid vector as described in U.S. Patent Application No. PCT/us02/41503, filed December 26, 2002. The entirety of this application is incorporated by reference, but for purposes of simplicity we have included particular passages which describe the chloroplast double vector. The description provide below will help in the understanding of the plasmids shown in Fig 9-28 of this Application.

Chloroplast transformation has been accomplished only in a few Solanaceous crops so far. There are several challenges in extending this technology to other crops. So far, only green chloroplasts have been transformed in which the leaf has been used as the explant. However, for many crops, including monocots, cultured non-green cells or other non-green plant parts are used as explants. These non-green tissues contain proplastids instead of chloroplasts, in which gene expression and gene regulation systems are quite different. During transformation, transformed proplastids should develop into mature chloroplasts and transformed cells should survive the selection process during all stages of development. Therefore, the major challenge is to provide chloroplasts the ability to survive selection in the light and the dark, at

different developmental stages. This is absolutely critical because only one or two chloroplasts are transformed in a plant cell and these plastids should have the ability to survive the selection pressure, multiply and establish themselves while all other untransformed plastids are eliminated in the selection process. The Double Barrel Plastid Vectors accomplish this by using genes coding for two different enzymes capable of detoxifying the same selectable marker (or spectrum of selectable markers), driven by regulatory signals that are functional in proplastids as well as in mature chloroplasts.

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The plastid vector described herein is one of several such non-limiting examples. The chloroplast flanking sequence contains appropriate coding sequences and a spacer region into which the transgene cassette is inserted. Any spacer sequence within the plastid genome could be targeted for transgene integration, including transcribed and transcriptionally silent spacer regions. Both aphA-6 and aphA-2 (nptII) genes code for enzymes that belong to the aminoglycoside phosphotransferase family but they originate from different prokaryotic organisms. Because of prokaryotic nature of the chloroplast genome, these genes are ideal for use in transgenic chloroplasts without any codon optimization. Genes of prokaryotic origin have been expressed at very high levels in transgenic chloroplasts (up to 47% of total soluble protein, DeCosa et al., 2001). Both enzymes have similar catalytic activity but the aphA-6 gene product has an extended ability to detoxify kanamycin and provides a wider spectrum of aminoglycoside detoxification, including amikacin. The advantage of choosing kanamycin as a selectable marker is that it has no natural resistance, unlike spectinomycin resistance observed in most monocots or spontaneous point mutation of the 16 S rRNA gene observed during the selection process. In addition, kanamycin is not in human clinical use as an antibiotic and several crops containing kanamycin resistant nuclear transgenes have been already approved by FDA for human consumption (e.g. flavor savor tomatoes) and currently in the market place.

As shown in Fig 36, in this non-limiting example, all transgenes are regulated by the plastid Prrn promoter; this 16S rRNA promoter drives the entire rRNA operon in the native chloroplast genome and contains binding sites for both the nuclear encoded and plastid encoded RNA polymerases. Therefore, this promoter is capable of functioning in both proplastids and chloroplasts (green and non-green, in the light and

The aphA-6 gene is further regulated by the gene 10 5' UTR capable of efficient translation in the dark, in proplastids present in non-green tissues (see GFP expression in proplastids of non-green cells of corn and carrot in Figs 32 and 2 regulated by the 16S rRNA promoter and gene 10 UTR). The rps16 3' UTR has been used to stabilize aphA-6 gene transcripts. The aphA-2 (nptII) gene, on the other hand is regulated by the psbA promoter, 5' and 3' UTRs, which are light regulated and highly efficient in the light, in chloroplasts (see A. Fernandez-San Millan, A. Mingeo-Castel, M. Miller and H. Daniell, 2003, A chloroplast transgenic approach to hyperexpress and purify Human Serum Albumin, a protein highly susceptible to proteolytic Plant Biotechnology Journal, in press; also see WO 01/72959). degradation. Therefore, a combination of both aphA-6 and aphA-2 genes, driven by regulatory signals in the light and in the dark in both proplastids and chloroplasts, provides continuous protection for transformed plastids/chloroplasts around the clock from the selectable agent. The gene(s) of interest with appropriate regulatory signals (gene X) are inserted downstream or upstream of the double barrel selectable system. Because multiple genes are inserted within spacer regions (DeCosa et al 2001, Daniell & Dhingra, 2002), the number of transgenes inserted does not pose problems in transcription, transcript processing or translation of operons (WO 01/64024). In a variation of this example, aphA-6 and aphA-2 genes, coupled with different transgenes are inserted at different spacer regions within the same chloroplast genome using appropriate flanking sequences and introduced via co-transformation of both vectors

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## Plant material transformation and regeneration of transgenic plants.

Sterile carrot plants (*Daucus carota* L. cv. Half long) were raised in plant tissue culture tubes containing MS salts (Murashige and Skoog, 1962), B5 vitamins (Gamborg et al 1968), 2% sucrose and 0.8% agar in the medium. The hypocotyls were cut into 0.5 mm segments and placed in 50 ml MSB liquid medium containing 3% sucrose, 0.1 mg/l 2,4-dichlorophenoxyacetic acid (2,4-D) and pH 5.7. After 3 weeks of continuous shaking at 26±2°C and 120 rpm, liberated cells were collected on a 100 μM mesh, centrifuged (150 x g for 10 min) and resuspended in fresh medium. Rapidly growing homogenous yellow cells were subcultured weekly. Fine cell suspension culture of carrot, filtered through a 100 μM mesh, were evenly spread on MSB solid medium supplemented with 3 mg/L 2,4-D and 1 mg/L kinetin and bombarded with

carrot specific plastid vectors (Fig. 1A-B). Bombarded calli incubated for 2 days in the dark were selected on MSB medium containing 3 mg/L 2,4-D, 1 mg/L kinetin and different concentrations of betaine aldehyde (10, 15, 20 and 25 mM BA) and spectinomycin (150, 250, 350 and 450 mg/L). Cultures were incubated in 16/8 h day/night cycle at 50-100 lx light intensity and 26±2°C temperature. Transgenic cultures were multiplied using both solid and liquid medium supplemented with selection agent. Transgenic plants produced through somatic embryogenesis (on MSB medium containing 0.2 mg/L) were transferred to soil in pots. The pots were covered with plastic bags to maintain high humidity for the first week and irrigated with progressively reduced concentrations of MS salts for the first week, followed by tap water in the second week.

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**DNA extraction, PCR and Southern blot analysis.** Total genomic DNA was isolated using a Plant DNeasy kit (Quiagen Inc. USA) for PCR and Southern blot analysis of transgenic carrot. PCR reactions were performed by denaturing 50 ng DNA template at 94°C for 5 min, running 25 PCR cycles (1min at 94°C, 1min at 64°C, 3min at 72°C), final extension at 72°C for 10 min, using Taq DNA polymerase with 10x PCR buffer and primers pairs 3P/3M (land on flanking sequence/land on *aadA* gene) and 16SF/1M (landing on the native chloroplast genome/ the *aadA* gene).

For Southern blot analysis, plant DNA was digested with PvuII and AflIII and transferred to nylon membranes for hybridizing with a 4.9 kb probe, generated by digesting *pDD-Dc-aadA/badh* vector DNA with PvuII and AflIII and labeled with <sup>32</sup>P using the ProbeQuant G-50 Columns (Amersham, USA). Blot was hybridized with probe using the Stratagene Quick-HYB hybridization solution and vender's instructions (Stratagene, USA).

BADH enzyme activity and immunoblot analysis. Extraction and assay for BADH (Betaine aldehyde dehydrogenase) activity was done as described (Daniell etal 2001). 1 g carrot tissues were grounded in 2 mL buffer containing 50 mM Hepes-KOH (pH 8.0), 1 mM EDTA, 20 mM Sodium metabisulfite, 10 mM Sodium borate, 5 mM ascorbic acid and 5 mM DTT. Crude extract was centrifuged at 10, 000 x g at 4°C for 10 min and the supernatant was desalted using Sephadex G-25 Columns (Amersham USA).  $NAD^{+}$ reduction by BADHwas measured biotech, Pharmacia spectrophotometrically at 340 nm after 1 min and 10 min in 1 mL assay buffer (50 mM

Hepes-KOH, pH 8.0; 1 mM EDTA, 5 mM DTT, 1 mM NAD<sup>+</sup>) added with 1 mM BA at 25°C to start reaction.

For immunoblot analysis total soluble protein was isolated using 2x Laemmli buffer from 100 mg carrot tissues. The mixture was boiled for 5 min and centrifuged for 5 min at 10,000 x g. Supernatant containing 50 µg total soluble protein (quantified with Bradford assay) was loaded on a 10% SDS-PAGE gel and transferred on blotting membrane. Membrane was hybridized with polyclonal anti-BADH serum, raised in rabbits against BADH (provided by Dr. Elisa Soto). Hybridized peptides were detected using horseradish peroxidase-linked secondary antibody, using Lumi-Phos<sup>TM</sup> WB chemiluminescent reagent (Pierce, USA).

Analysis of transgenic plants for salt tolerance. Transgenic and non-transgenic carrot plants transferred to soil in pots of same morphogenic growth phase and height were analyzed for salt tolerance containing 0, 100, 200, 300, 400 and 500 mM NaCl, respectively. Plants were maintained in growth chamber and irrigated daily with saline water containing different levels of salt for one month.

# ILLUSTRATIVE EXAMPLE 2 COTTON TRANSFORMATION

Material and Methods

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Plant material and transformation: Delinted cotton (Gossypium hirsutum L. cv. Coker310FR) seeds were sterilized by dipping in 70% ethanol for 2 minutes followed an 8 minutes treatment with sodium hypochlorite solution containing approximately 4% available chlorine and then by treatment with 0.1% mercuric chloride solution (w/v) for 5 minutes. After surface sterilization and four to five washes with sterile water, seeds were kept in sterile water for 4-5 hours for softening the seed coat which was completely removed before the seeds were placed on 1/2 MSB medium containing half strength MS salts (Murashige and Skoog, 1962) and B5 vitamins (Gamborg etal 1968) with 1.5% sucrose. Hypocotyl explants (4-6 mm long) of 5 day old seedlings were placed vertically on MST1 medium (containing MS salts, B5 vitamins, 0.1 mg/l 2,4-D, 0.5 mg/l kinetin and 3% glucose) for the induction of callus. Uniformly distributed pro-embryognic callus were bombarded with gold particle coated with chloroplast vector using the Helium-based biolistics particle gun (Bio-Rad). The transformation of cotton callus was optimized using the parameters: 0.6μm gold

particles macro-carrier; 27 in. Hg chamber vacuum; 1550 psi Helium pressure; 6 mm rupture disc macro-carrier gap; 6 mm macro-carrier flying distance, 6 cm target distance and 10 µl chloroplast vector coated on gold particles. Cultures after bombardment were incubated in dark for 24 h and thereafter transferred to selection medium MST1 supplemented with 50 mg/l kanamycin and incubated in 16/8 h day/night cycle at 750 lux light intensity and 28±2°C temperature. Transgenic embryogenic cultures were multiplied on MST1 medium supplemented with 50 mg/l kanamycin (Fig 34) and transgene aphA-6 gene integration in cultures was confirmed by PCR (Fig 35).

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# ILLUSTRATIVE EXAMPLE 3 EXPRESSION CASSETTE CONSTRUCTION

#### Materials and Methods:

Amplification and cloning of flanking sequences: DNA fragment representing flanking sequences were amplified from plant genomic DNA that was isolated from the leaves using Qiagen plant extraction kit following manufacturer's protocol. The flanking sequence fragment was amplified with the primers, ADLF – 5' gtgtcagtgtcggccagcagag 3' and ADLR – 5' aacaggggtcaaggtcggccag 3' using Platinum Pfx DNA polymerase (Invitrogen Inc.). The amplified fragment represents the 16S/trnI-trnA/23S region of the chloroplast genome and is approximately 4.2 kb in size. The PCR amplified DNA fragment was treated with T4 polynucleotide kinase (Promega) and cloned into PvuII digested pBluescript II KS dephosphorylated with Shrimp Alakaline phosophatase (Promega). The kinase and dephosphorylation reactions were performed as per the manufacturer's instructions. The clone harboring carrot specific flanking region was designated as pDA-35.

Construction of the expression cassettes: pDA-29 is a chloroplast specific expression cassette cloned in pBluescript II KS that carries the *aadA* gene conferring resistance to spectinomycin and streptomycin and *badh* gene that metabolizes the breakdown of toxic betaine aldehyde to glycinebetaine. Expression of the first gene is driven by the 16S Prrn promoter, under the regulation of a Shine-Dalgarno sequence at the 5' end and *psbA* 3' UTR at the 3' end. Expression of *badh* gene is regulated by heterologous T7 gene 10 5'UTR and rps16 3'UTR. The *aadA* gene was derived from

pLD-CtV (Daniell et al. 1998) and the *badh* gene was derived from pLD-BADH (Daniell et al. 2001). Second chloroplast expression cassette pDA-30 carries the *sm-gfp* gene encoding for soluble modified green fluorescent protein (obtained from TAIR) and the *badh* gene. Expression of *sm-gfp* is driven by the 16S Prrn promoter and is regulated by heterologous T7 gene 10 5'UTR and rps16 3'UTR. The expression of *badh* gene is regulated by *psb*A promoter/5' UTR and 3' UTR. The promoters and regulatory sequences were amplified using PCR based on the information available for tobacco chloroplast genome (Shinozaki et al. 1987).

Construction of chloroplast transformation vectors: Chloroplast transformation vector pDD-XX-aadA/badh is a derivative of pDA vectors that harbor the flanking sequences with the pDA-29 expression cassette. The expression cassette from pDA-29 was obtained as an ApaI fragment, blunt-ended using klenow DNA polymerase (NEB) as per manufacturer's instructions and cloned into PvuII digested and dephosphorylated pDA-35. The other species specific chloroplast transformation vector pDD-XX-gfp/badh harbors the carrot flanking sequence and the pDA-30 expression cassette. The expression cassette from pDA-30 was derived as a ClaI/SacI fragment, blunt-ended and cloned into PvuII digested and dephosphorylated pDA-35. Bacterial and DNA manipulations were performed as per standard molecular biology protocols.

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# ILLUSTRATIVE EXAMPLE 4: CORN TRANSFORMATION

For genetic engineering of the corn chloroplast genome, corn specific sequences, flanking the targeted integration site in the corn chloroplast genome (trnI and trnA) were amplified with specific PCR primers and subcloned to flank the betaine aldehyde dehydrogenase (BADH) selectable marker, and green fluorescent protein (GFP) reporter gene expression cassette.

Callus cultures were initiated from aseptically excised immature zygotic embryos (1-2 mm in length), produced on self-pollinated ears of HiII (F1) maize plants. Ears were surface sterilized in a solution containing 2.6% Sodium hypochlorite (prepared with commercial bleach) containing .1% Tween 20 (polyoxyethylene sorbitan monolaurate) for 20 miniutes under continuous shaking, then rinsed 4 timesin sterile distilled water. The Embryos were then placed on the callus induction medium

CI-1, which contained N6 salts and vitamins (463.0mg/l (NH4)2SO4, 2830.0mg/lKNO3, 400mg/l KH2PO4, 166.0 mg/l CaCl2, 185 mg/l MgSO4. 7H2O, 37.3 mg/l Na2-EDTA, 27.85mg/l FeSO4.7H2O, 1.6 mg/l H3BO3, 4.4 mg/l MnSO4.H2O, .8, KI, 1.5mg/l ZnSO4. 7H2O), 2% sucrose and 1.0 mg/l 2,4-D (2,4 dichloro-phenoxy acetic acid), with the rounded scutellar side exposed and the flat plumule-radicle axis side in contact with the medium. Callus cultures were maintained in darkness at 25-28 °C and subcultured every two weeks.

Particle bombardment of embryogenic calli

Micro projectiles were coated with DNA (pDA34-ZM-gfp-BADH and pDA33-ZM-aadA-BADH) and bombardment was carried out with the biolistic device PDS1000/He (Bio-Rad).

Prior to bombardment, embryogenic calli were selected, transferred over sterile filter paper (Watman No.1), and placed on the surface of a fresh medium in standard Petrti plates (100x 15mm). Gold particles (0.6μm) were then coated with plasmid DNA as follows: 50μl of washed gold particles were mixed with 10μl DNA (1μg/μl), 50μl of 2.5M CaCl2, 20μl of 0.1M spermidine and vortexed. Particles were cneterfuged for a few seconds at 3000rpm and then the ethanol was poured off. Ethanol washing was repeated five times, then the pellet was resuspended in 30μl of 100% ethanol and placed on ice until it was used for bombardment (the coated particles were used within 2hours). Bombardment was carried out with the biolistic device PDS1000/He (Bio Rad) by loading the target sample at level 2 in the sample chamber under a partial vacuum (28 inches Hg).

The callus cultures were bombarded with the maize chloroplast transformation vectors using 1100 psi rupture discs. Following bombardment, the explants were transferred to a fresh medium; plates were sealed with micropore tape and incubated in darkness at 25-28 °C.

#### Selection

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Selection was intiated two days after bombardment. The bombarded calli were transferred to callus induction medium containing 5-20mM BA (betaine aldehyde) or 25-100mg/l streptomycin. Selection was also carried out using 50-150 mM NaCl in combination with the BA to maintain osmostic pressure.

## Regeneration

Regeneration was initiated 6 to 8 weeks after bombardment by transferring the calli to a medium R1 containing Ms salts and vitamins supplemented with 1.0 mg/l NAA ( $\alpha$ -naphthalene acetic acid),2% sucrose, 2g/l myoinositol and .3% phytagel at pH 5.8. Regenerated plants were transferred to R2 containing ½ MS salts and vitamins, 3% sucrose and .3% phytagel at pH 5.8. Regenerated plants were maintained in light (16 /8 hr photoperiod).

Shoot multiplication

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The surface steriliztion and germination of corn seeds

Corn seeds were surface sterilized in a solution containing 2.6% Sodium hypochlorite (prepared from commercial bleach) containing .1% Tween 20 for 20 minutes under continuos shaking, then rinsed four times in sterile distilled water. Seeds were grown on MS medium at pH 5.8 in darkness. Nodal sections were excised aseptically from three day old seedlings. The nodal sections appear as clear demarcations on the germinated seedlings and represent the seventh node. When excised, the nodal cross sections are 1.3 to 1.5 mm in length.

Particle bombardment of nodal sections

Prior to bombardment, 20-30 nodal sections were placed in the center of each petri plate with acropitila end up. Bombardment was carried out with the maize chloroplast vectors, using 1100, 1300 and 1550 psi rupture discs.

Multiple shoot induction and selection

Nodal section explants are placed acropital end up on shoot multiplication medium SM1 composed of Ms salts and vitamins, 1.0 mg/l 6BA (6-Benzyl amino purine), 3% sucrose and 5g/l phytagel at pH 5.8 under continuous light at 25°C. Initiation of the shoot-tip clumps from the original shoot tips occurred 2 to 4 weeks after culture. Two days after bombardment, transformed nodal sections were transferred to shoot multiplicaton medium containing 5-20mM BA or 50-100 mg/l streptomycin selective agents. Subsequent subcultures at two week intervals were carried out by selecting, dividing and subculturing green clumps on selective shoot multiplication medium containing 5-20mM BA or 25-100 mg/l streptomycin.

Regeneration

The Multiple shoot clumps were regenerated by transferring them to regeneration medium M1 containing MS salts and Vitamins, 5 mg/l IBA and 3%

sucrose at pH 5.8. The developed shoots were regenerated by transferring the shoot tip clumps to M2 medium containing ½ MS salts and vitamins, 3 % sucrose and 3g/l phytagel at pH 5.8. It should be further rnoted that all the regeneration media are supplemented with 5-20mM BA or 25-100mg/l streptomycin as the selective agents.

To engineer the corn chloroplast genome free of antibiotic resistance genes, maize calli were bombarded with a chloroplast expression vector containing the green fluorescent protein (GFP) and the betaine adehyde dehydrogenase (BADH) genes as selectable or screenable markers. To compare the betaine aldehyde (BA) selection with streptomycin, another chloroplast expression vector was constructed containing the aadA and the BADH genes. The number of putative transgenic events was higher on BA selection than on streptomycin. Transgenic corn tissues screened on BA were examined using a laser-scanning confocal microscope. The GFP fluorescence was observed throughout the somatic embryos of corn. Chloroplast transformation of corn provides a suitable avenue for the production of edible vaccines and oral delivery of biopharmaceuticals.

Corn chloroplast transformation vector

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Corn chloroplast transformation vector facilitates the integration of transgene into the inverted repeat (IR) region of the corn chloroplast genome. The vector pLD-Corn-BADH contains the chimeric aadA gene and the BADH gene driven by the constitutive 16 S rRNA promoter and regulated by the 3' UTR region of psbA gene from petunia plastid genome. In this construct both, aadA and BADH possess the chloroplast preferred ribosomal binding site, GGAGG. Another vector used for corn chloroplast transformation pLD-corn-UTR-BADH has the constitutive 16 S rRNA promoter driving the expression of the dicistron, but BADH is under the regulation of the promoter and the 5' UTR of the psbA gene and the 3' UTR of psbA gene, for Since the expression of the foreign protein is desired in enhanced expression. chromoplasts of corn seeds, the gene of interest needs to be under the control of a regulatory sequence that is free from cellular control. In this context, examples of suitable candidate regulatory sequences are the T7 gene 10-leader sequence and cry2Aa2 UTR. The T7 gene 10-leader sequence is used to express foreign proteins in transgenic chromoplasts. The cry2Aa2 UTR has been shown by the inventor to accumulate as much foreign protein in chromoplasts as efficient as the psbA UTR in

green tissues. Therefore the selectable marker for additional vectors use the BADH gene under the regulation of psbA promoter and 5'UTR, as psbA is one of the most efficiently translated chloroplast genes in green tissues. When green tissue or non-green embryogenic calli are used for introducing the transgene into the corn chloroplast genome, it is preferred to use the light regulated psbA promoter/ UTR or 16 S rRNA promoter/gene 10 UTR, respectively.

The following list of examples wherein the Applicant has demonstrated somatic embryogenesis in corn, cotton, and carrot clearly demonstrate that the transformation of any of a number of plants is suitable owing to the Applicants description and examples contained herein. The ability to transform any of a number of plants so that these plants perform embryogenesis is further bolstered by the number of exemplary references listed below which illustrate a methodology of plant transformation through somatic embryogenesis in a wide variety of plants. The following references are examples of regeneration of crops via embryogenesis and/or a crop/plant transformation via the nuclear genome. As a result, one skilled in the art can use the the protocols for somatic embryogenesis described in the exemplary references below, in concert with the chloroplast vectors described in the specification, to achieve plastid transformation through non-green plant cells. In other word, one skilled the art will recognize that the protocols below are applicable to the description in the content of the specification. This following list of references, hereby fully incorporated into this application, describe plant transformation through somatic embryogenesis:

# **Crops:**

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Maize:

J. F. Petolino, N. L. Hopkins, B. D. Kosegi, M. Skokut genetic transformation and hybridization: Whisker-mediated transformation of embryogenic callus of maize. 19: 781-786 (2000).

Soybean:

H. N. Trick, J. J. Finer. Sonication-assisted Agrobacterium-mediated transformation of soybean [Glycine max (L.) Merrill] embryogenic suspension culture tissue. Plant Cell Reports 17: 482-488 (1998).

Wheat:

A. Pellegrineschi, R. M. Brito, L. Velazquez, L. M. Noguera, W. Pfeiffer, S. McLean, D. Hoisington. The effect of pretreatment with mild heat and drought stresses on the explant and biolistic transformation frequency of three durum wheat cultivars. Plant Cell Rep 20: 955-960 (2002).

Barley:

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M. Manoharan, L. S. Dahleen. Genetic transformation of the commercial barley (Hordeum vulgare L.) cultivar Conlon by particle bombardment of callus

Plant Cell Rep 21: 76-80 (2002).

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## **CLAIMS**

- 1. A plastid transformation vector suitable for transforming a non-green plant cell, said plastid vector comprising, as operably linked components, a first flanking sequence, a DNA sequence coding for a foreign gene, and a second flanking sequence.
- 2. The vector of Claim 1, further comprising a regulatory sequence functional in said plastid transformation vector.

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- 3. The vector of Claim 2, wherein said regulatory sequence comprises a promoter operative in said plastid genome.
  - 4. The vector of Claim 3, wherein said promoter is *Prrn 16S* rRNA.
- 5. The vector of Claim 4, wherein said regulatory sequence comprises psbA 5' and psbA 3' elements.
- 6. The vector of Claim 2, wherein said regulatory sequences further comprise a 5' UTR capable of providing transcription and translation enhancement of said DNA sequence coding for a foreign gene.
- 7. The vector of Claim 2, wherein said regulatory sequences further comprise a 3' untranslated region (UTR) capable of conferring transcript stability to said DNA sequence coding for a foreign gene.
- 8. The vector of Claim 2, wherein said regulatory sequence further 20 comprise a gene 10 5'UTR.
  - 9. The vector of Claim 8, wherein said regualtory sequence further comprises a gene 10 5'UTR/rps 16 3'UTR.
  - 10. The vector of Claim 1, wherein said first flanking sequence is 16S/trnI, and wherein said second flanking sequence is trnA/23S.
    - 11. The vector of claim 1, wherein said first flanking sequence is about 4kb.
  - 12. The vector of claim 1, wherein said second flanking sequence is about 4kb.
  - 13. The vector of Claim 1, wherein the vector is component for stably integrating into a plastid genome of a higher plant species, and wherein said first and second flanking DNA sequences are substantially homologous to sequences in a spacer region of said plastid genome, and wherein said flanking sequences are conserved in the plastid genome of said higher plant species.

14. The vector of Claim 13, wherein said spacer region is a transcriptionally active spacer region.

- 15. The vector of Claim 6, wherein said 5' UTR is a 5' UTR of psbA.
- 16. The vector of Claim 7, wherein said 3' UTR is a 3' UTR of psbA.
- 5 17. The vector of Claim 1 further comprising a DNA sequence encoding a selectable marker.
  - 18. The vector of Claim 17, wherein said selectable marker is an antibiotic-free selectable marker.
- 19. The vector of Claim 18, wherein said antibiotic-free selectable marker is 10 Betaine aldehyde dehydrogenase (BADH).
  - 20. The vector of Claim 17, wherein said DNA sequence encoding said selectable marker encodes an antibiotic resistance selectable marker.
  - 21. The vector of Claim 20, wherein said antibiotic resistance selectable marker is *aadA*.
  - 22. A plant stably transformed with the vector of Claim 1.
    - A progeny of the plant of claim 22.
    - 24. A seed of the plant of claim 22.

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- 25. A non-green part of the plant of claim 22, comprising a plastid genome having a heterologous DNA sequence coding for polypeptide of interest.
- 26. The plant of Claim 22, wherein said plant further comprises at least one pro-plastid transformed with the vector of Claim 1.
  - 27. A somatic embryo transformed with the vector of Claim 1.
  - 28. A non-green plant cell comprising a non green plastid including an expression cassette comprising, as operably joined components, a promoter functional in said non-green plastid, heterologous DNA sequence encoding a polypeptide of interest, at least one additional structural gene or functional portion thereof encoding a polypeptide which confers a selectable trait, wherein transcription of said heterologous DNA sequence is regulated by said promoter, a transcriptional termination region and

plastid DNA flanking sequences flanking said expression cassette to facilitate stable integration of the expression cassette into a genome of said plastid by homologous recombination.

29. The non-green plant cell of Claim 28, capable of regenerating through somatic embryogeneis.

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- 30. A transgenic non-green plant cell having a plastid genome transformed with the plastid transformation vector of Claim 1, wherein said transgenic non-green plant cell is regenerated through somatic embryogenesis.
- 31. A method of transforming a plastid through somatic embryogenesis comprising the steps of:

integrating the vector of Claim 1 into a plastid genome of a plant plastid.

- 32. A method of achieving plastid transformation using no-green explants, wherein a plant is regenerated through somatic embryogenesis comprising the steps of:
- 33. A plant cell comprising a plastid including an expression cassette comprsing, as operably joined components, a heterologous DNA sequence encoding a polypeptide of interest, a DNA sequence encoding a selectable marker, and plastd DNA sequences flanking the expression cassette to facilitate stable integration of the said expression cassette into the chloroplast genome by homologous recombination, wherein said plant cell is regenerated through somatic embyogenesis.
- 34. A non-green plant cell capable of regenerating through somatic embryogenesis, wherein said non-green plant cell comprises a heterologous DNA sequence coding for a polypeptide of interest, wherein said heterologous DNA sequence coding for said polypeptide of interest is integrated into a plastid contained in said non-green plant cell.
- 35. A plastid transformation vector capable for transforming non-green plant cells, said plastid vector comprising, as operably linked components, a first flanking sequence, promoter operative in a plastid, a DNA sequence coding for a selectable marker operative in said plastid, a DNA sequence coding for a foreign gene, and a second flanking sequence.
- 36. A plastid transformation vector suitable for transforming a plastid, wherein said plastid to be transformed is in a non-green plant cell, and wherein said plastid transformation vector comprises, as operably linked components, a first

flanking sequence, a regulartory sequence functional in said plastid, a heterologous DNA sequence coding for a polypeptide of interest, and a second flanking sequence.

37. A method for producing a polypeptide of interest in a non-green plant cell, wherein said polypeptide of interest is coded for by a heterologous DNA sequence, comprising the steps of:

integrating a plastid transformation vector according to Claim 1 into the plastid genome of a plant cell; and

growing said plant cell to express said polypeptide of interest.

5

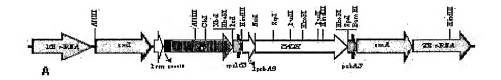
15

10 38. A method of visually selecting a transgenic plant comprising the steps of:

transforming a non-green plant cell via the vector of claim to express an exogenous betain aldhyde dehydrogenase (badh) gene.

39. A plastid transformation vector suitable for transforming a non-green plant cell, said plastid vector comprising, as operably linked components, a first flanking sequence, a regulatory sequence functional in a plastid, DNA sequence coding for a foreign gene, and a second flanking sequence.

FIG 1



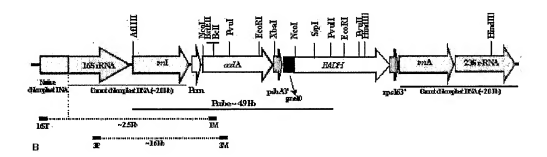


FIG 2

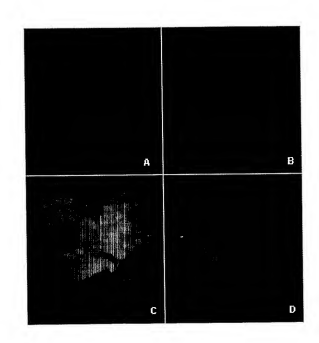


FIG 3

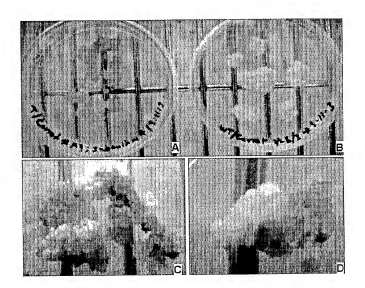
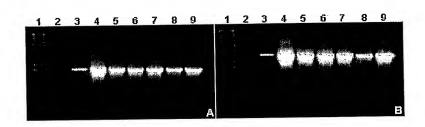


FIG 4



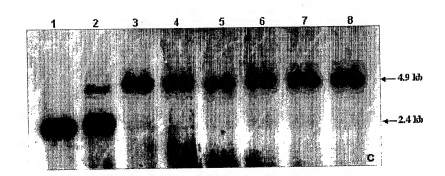
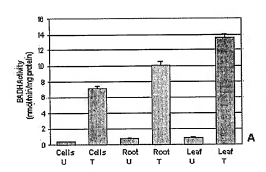


FIG 5



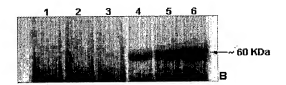


FIG 6

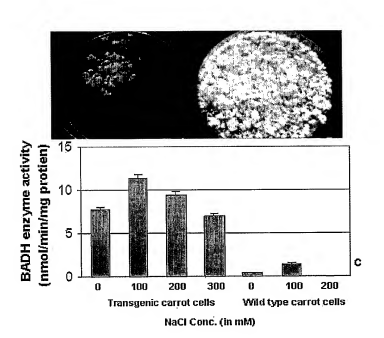


FIG 7

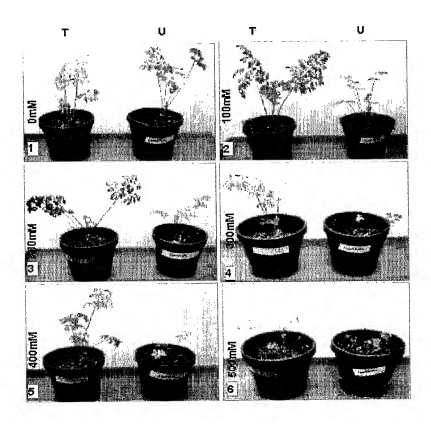
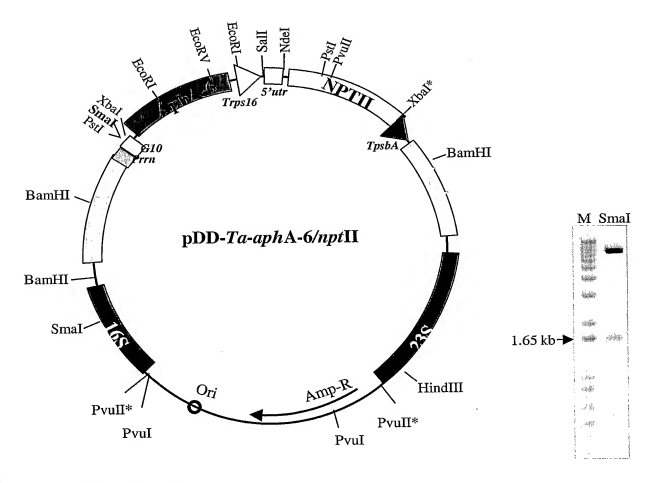


FIG8
PLASMID NAME: pDD-Ta-aphA-6/nptII



<sup>\*</sup> Means destroyed

FIG 9

PLASMID NAME: pDD-So-aphA-6/nptII

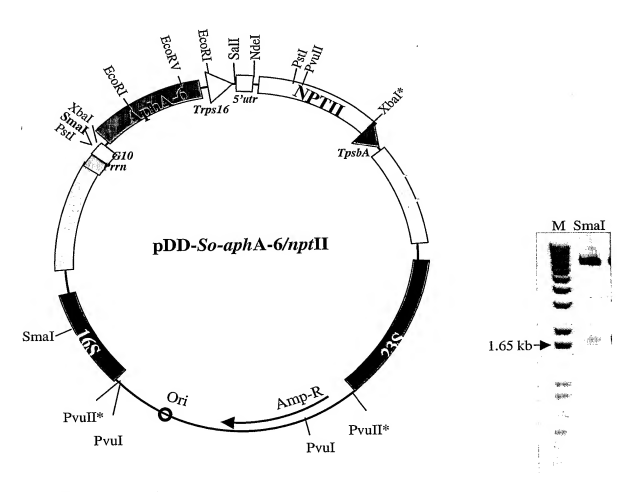


FIG 10 PLASMID NAME: pDD-Dc-aphA-6/nptII

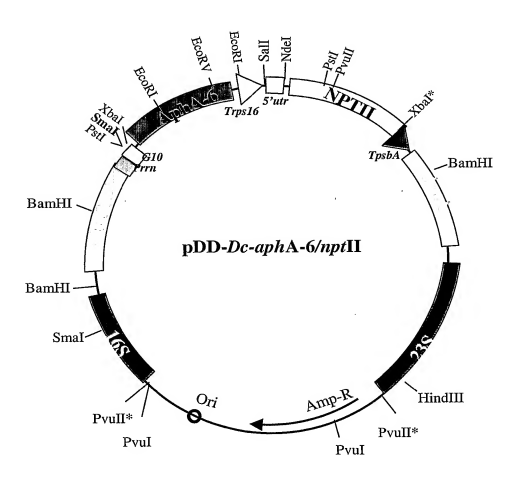
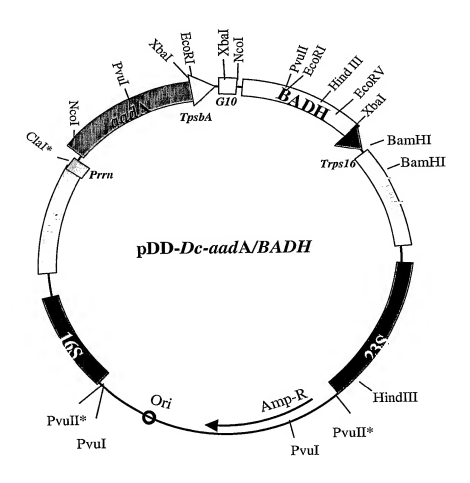


FIG 11
PLASMID NAME: pDD-Dc-aadA/BADH



<sup>\*</sup> Means destroyed

FIG 12
PLASMID NAME: pDD-Dc-gfp/BADH

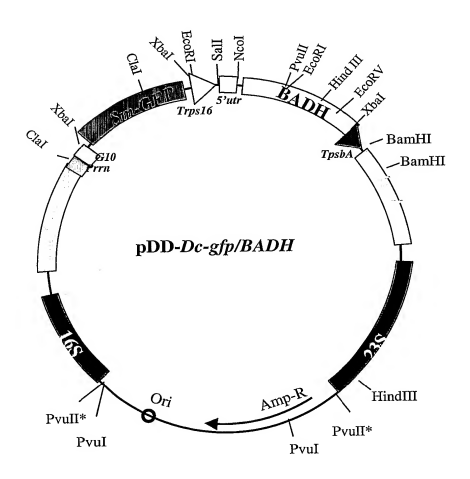


FIG 13
PLASMID NAME: pDD-Gh-aphA-6/nptII

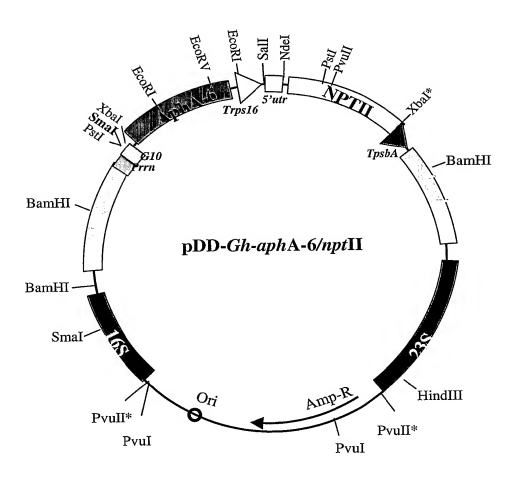
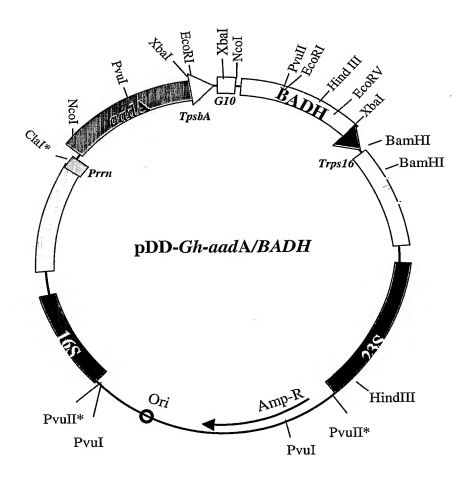
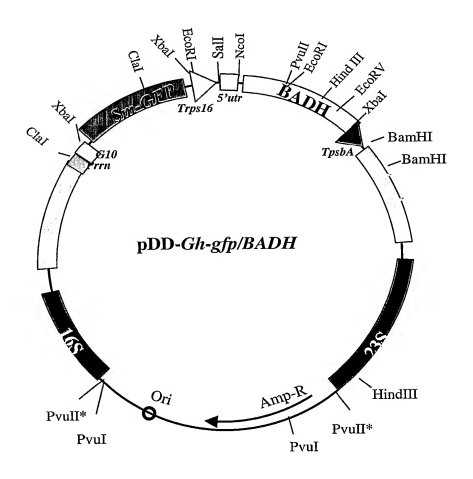


FIG 14
PLASMID NAME: pDD-Gh-aadA/BADH

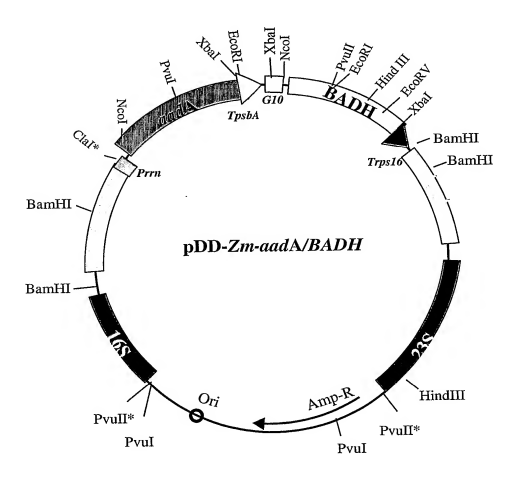


<sup>\*</sup> Means destroyed

FIG 15
PLASMID NAME: pDD-Gh-gfp/BADH



**FIG 16**PLASMID NAME: pDD-*Zm-aad*A/*BADH* 



**FIG 17**PLASMID NAME: pDD-Zm-gfp/BADH

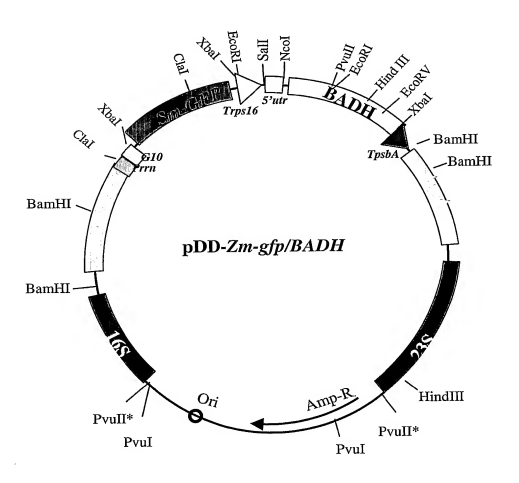


FIG 18

PLASMID NAME: pDD-Zm-aphA-6/nptII

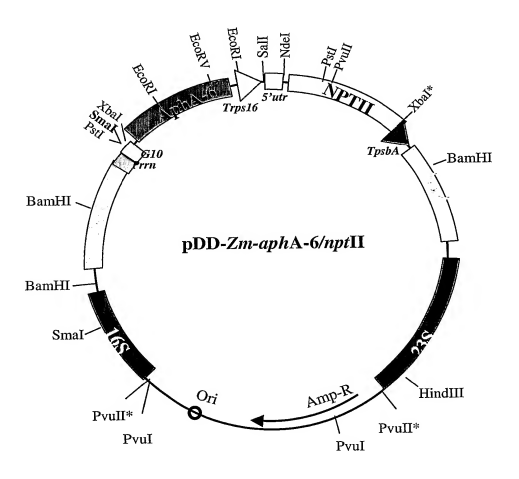


FIG 19
PLASMID NAME: pDD-Pv-aphA-6/nptII (switchgrass)

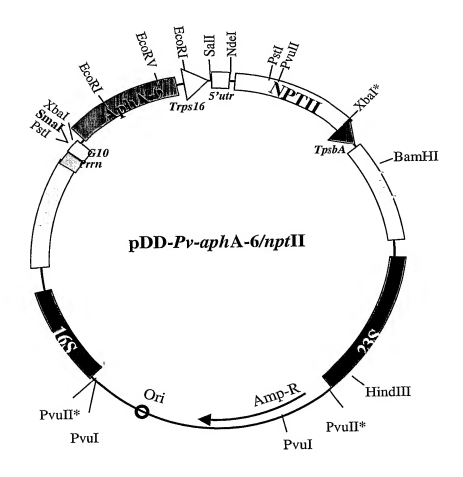
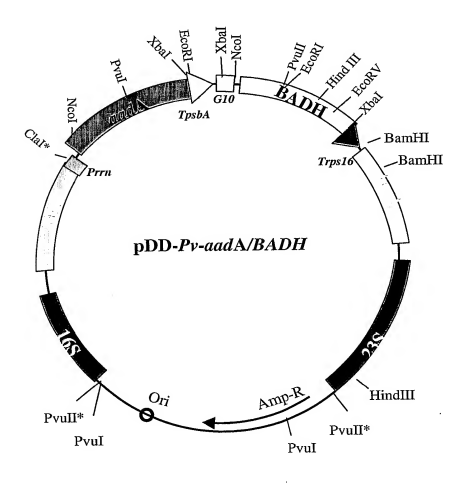


FIG 20
PLASMID NAME: pDD-Pv-aadA/BADH (switchgrass)



<sup>\*</sup> Means destroyed

FIG 21

PLASMID NAME: pDD-Cd-aphA-6/nptII (bermudagrass)

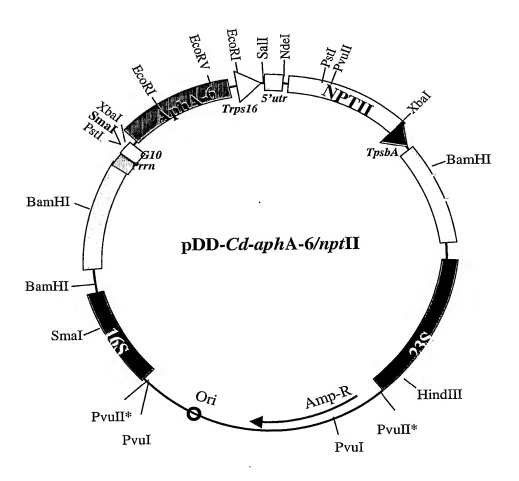
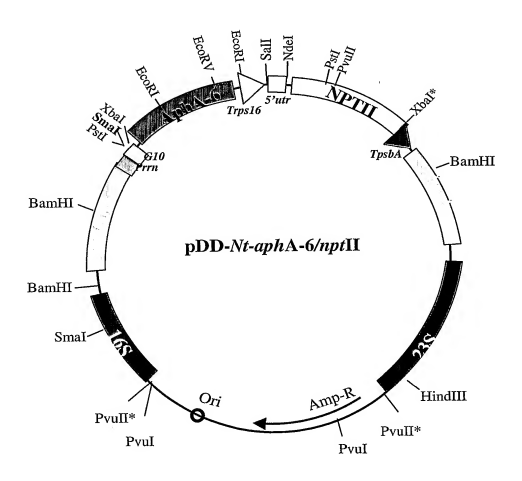
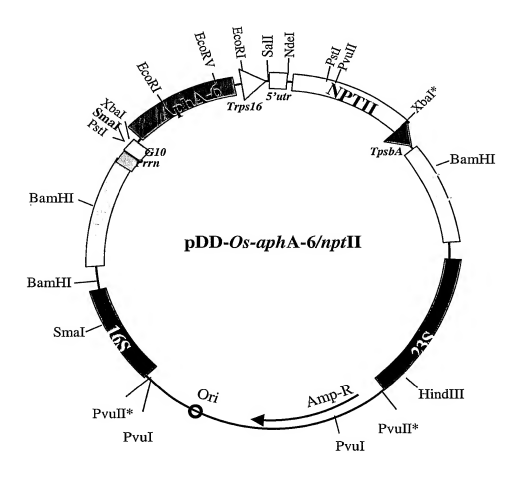


FIG 22
PLASMID NAME: pDD-Nt-aphA-6/nptII



<sup>\*</sup> Means destroyed

FIG 23
PLASMID NAME: pDD-Os-aphA-6/nptII



<sup>\*</sup> Means destroyed

FIG 24
PLASMID NAME: pDA-66

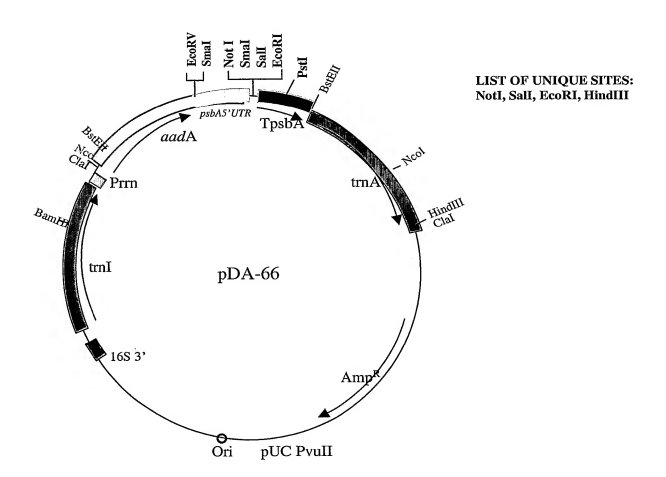
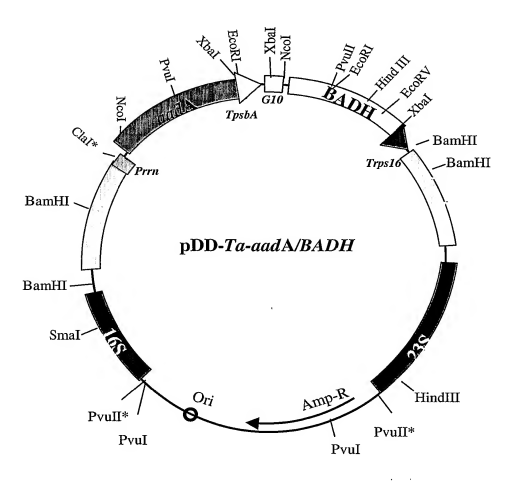


FIG 25
PLASMID NAME: pDD-*Ta-aadA/BADH* 



**FIG 26**PLASMID NAME: pDD-*Ta-gfp/BADH* 

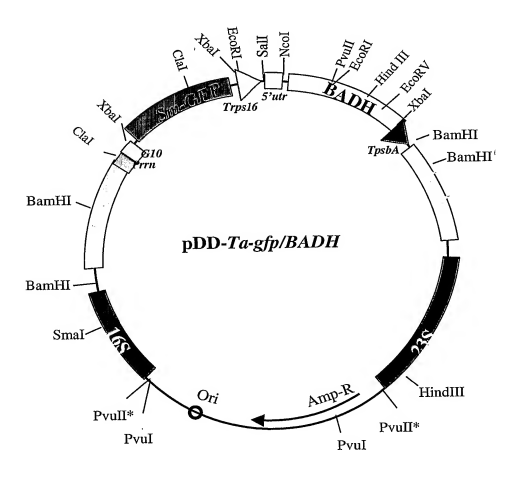
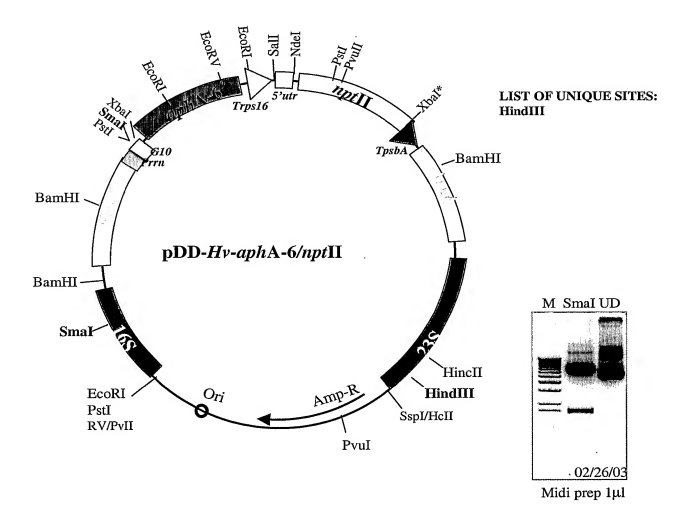
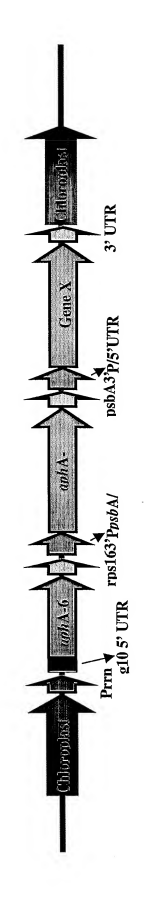


FIG 27
PLASMID NAME: pDD-Hv-aphA-6/nptII

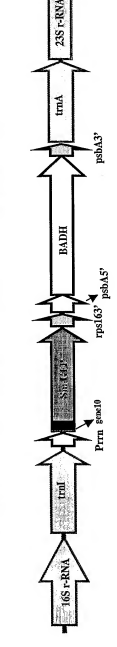




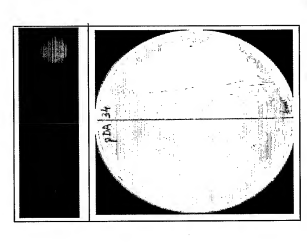
Double Barrel Plastid Vector harboring aphA-6 and aphA-2 genes conferring resistance to aminoglycosides

**FIG 28** 



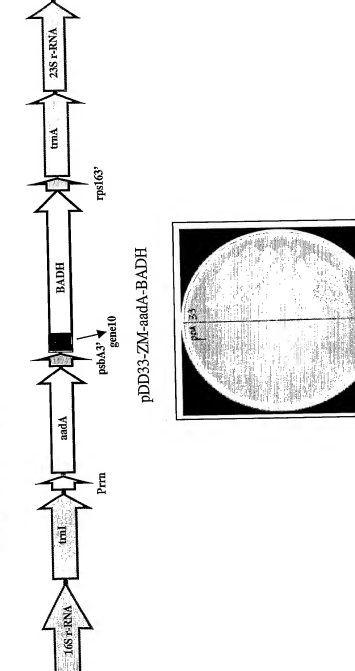


pDD34-ZM-gfp-BADH

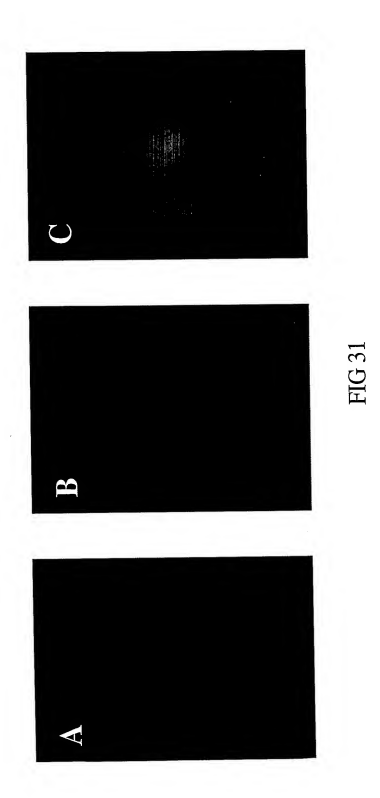


GFP expression in E. coli





E. coli cells grown on Spectinomycin



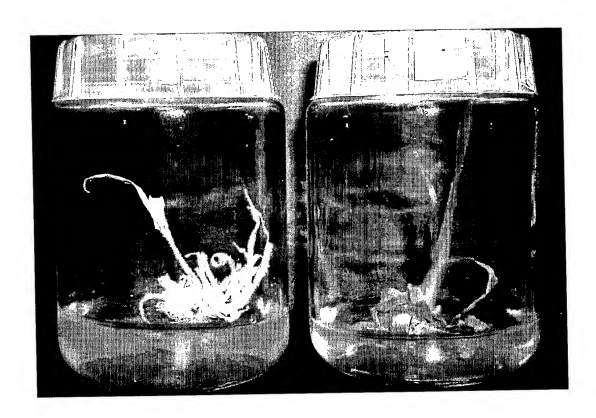


FIG 32A

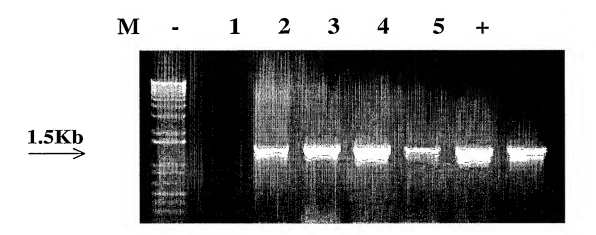


FIG 32B

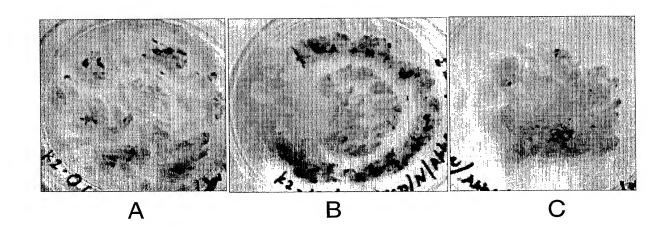
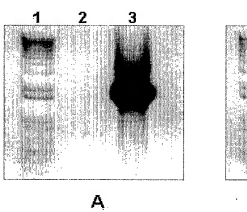
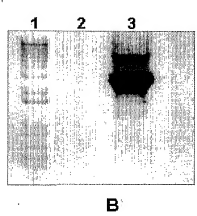


FIG 33(A-B)





Primers: 3P-aphA6 Primers: 16SF-aphA6

FIG 34 (A-B)

## **FIG 35**

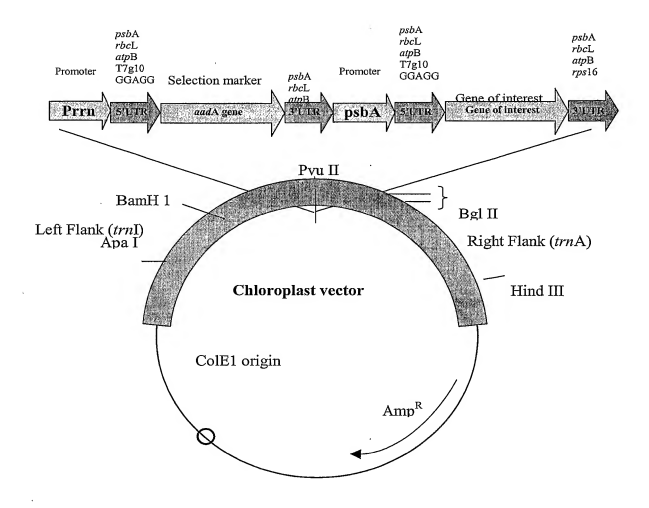
1. Sequence of aadA/BADH cassette (SEQ ID No. 1): AGCTTGCGGGCCCCCCTCGAGGTCGACGGTATCGATGAGCCTGATTATCCCTAAGCCCAATGTGAGTTTTTCTAGTTGG ATTTGCTCCCCCGCCGTCGTTCAATGAGAATGGATAAGAGGCTCGTGGGATTGACGTGAGGGGGCAGGGATGGCTATATT TCTGGGAGCGAACTCCGGGCGAATATGAAGCGCATGGATACAAGTTATGCCTTGGAATGAAAGACAATTCCGAATCCGCT TTGTCTACCCGATACAAGTGAGTTGTAGGGAGGCAACCATGGCAGAAGCGGTGATCGCCGAAGTATCGACTCAACTATCA GAGGTAGTTGGCGTCATCGAGCGCCATCTCGAACCGACGTTGCTGGCCGTACATTTGTACGGCTCCGCAGTGGATGGCGG CCTGAAGCCACACTGATATTGATTTGCTGGTTACGGTGACGGTGACCGTAAGGCTTGATGAAAACAACGCGGCGAGCTT TGATCAACGACCTTTTGGAAACTTCGGCTTCCCCTGGAGAGAGCGAGATTCTCCGCGCTGTAGAAGTCACCATTGTTGTG CACGACGACATCATTCCGTGGCGTTATCCAGCTAAGCGCGAACTGCAATTTGGAGAATGGCAGCGCAATGACATTCTTGC AGGTATCTTCGAGCCAGCCACGATCGACATTGATCTGGCTATCTTGCTGGCAAAAGCAAGAGAACATAGCGTTGCCTTGG TAGGTCCAGCGGCGGAGGAACTCTTTGATCCGGTTCCTGAACAGGATCTATTTGAGGCGCTAAATGAAACCTTAACGCTA TGGAACTCGCCGCCGACTGGGCTGCCGATGAGCGAAATGTAGTGCTTACGTTGTCCCGCATTTGGTACAGCGCAGTAACCGGCAGAATCGCCGCCGAAGGATGTCGCTGCCGACTGGCAATGGAGCGCCTGCCGGCCCAGTATCAGCCCGTCATACTTG AAGCTAGACAGGCTTATCTTGGACAAGAAGAAGATCGCTTGGCCTCGCGCGCAGATCAGTTGGAAGAATTTGTTCACTAC GTGAAAGGCGAGATCACCAAGGTAGTCGGCAAATAAAAAGCCGAATCTAGAGCGATCCTGGCCTAGTCTATAGGAGGTTT TGATTGAGTATTCTATTTTGATTTTGTATTTGGGCTGCGCGGGGAGACCACAACGGTTTCCCTCTAGAAATAATTT TGTTTAACTTTAAGAAGGAGATATACCATGGCGTTCCCAATTCCTGCTCGTCAGCTATTCATCGACGGAGAGTGGAGAGA ACCCATTAAAAAAAATCGCATACCCGTCATCAATCCGTCCACTGAAGAAATCATCGGTGATATTCCGGCAGCCACGGCTG AAGATGTGGAGGTTGCGGTGGCAGCTCGAAGAGCCTTTAGGAGGAACAATTGGTCAGCAACATCTGGGGCTCATCGT GAAACCTTTTGATGAAGCAGTGCTGGACATTGATGACGTTGCTTCATGTTTTGAATATTTTGCCGGACAAGCAGAAGCTC TTGATGGTAAACAAAAGGCTCCAGTCACCCTGCCTATGGAAAGGTTCAAAAGTCATGTTCTCAGGCAGCCCCTTGGTGTT GTTGGATTAATATCCCCATGGAATTACCCACTTCTAATGGCTACATGGAAAATTGCTCCAGCACTTGCTGCTGGGTGTAC AGCTGTACTTAAGCCATCCGAGTTGGCATCTGTGACTTGTCTAGAATTCGGTGAAGTTTGCAACGAAGTGGGACTTCCTC CAGGCGTGTTGAATATCTTGACAGGATTAGGTCCAGATGCTGGTGCACCATTAGTATCACACCCCGATGTTGACAAGATT GCCTTTACTGGGAGTAGTGCCACTGGAAGCAAGGTTATGGCTTCTGCTGCCCAATTGGTTAAGCCTGTTACATTAGAACT TGGGGGTAAAAGTCCTATTGTAGTGTTTGAAGATGTTGATATTGATAAAGTTGTGGAATGGACTATTTTTGGCTGTTTCT GGACAAATGGTCAAATATGTAGTGCAACGTCTAGACTGCTTGTGCATGAAAGTATTGCAGCTGAGTTTTGATAAGCTT GTAAAATGGACGAAAAACATTAAAATTTCTGACCCATTTGAAGAAGGATGCCGGCTTGGCCCTGTTATTAGTAAAGGACA GTACGACAAAATTATGAAGTTCATATCAACAGCAAAGAGTGAGGGGGCAACTATTTTGTATGGAGGTTCCCGTCCTGAGC ATTTGAAGAAAGGTTATTACATTGAACCCACCATTGTAACTGATATCTCCACATCCATGCAAATATGGAAAGAGGAAGTT TTTGGCCCTGTCTTGTGTGTTAAAACATTTAGTTCCGAAGATGAAGCCATTGCATTGGCAAATGATACAGAGTACGGTTT AGCTGCTGCTGTTTTCTAATGATCTTGAAAGATGTGAGAGGATAACGAAGGCTCTAGAAGTTGGAGCTGTTTGGGTTA ATTGCTCACAACCATGCTTTGTTCAAGCTCCTTGGGGAGGCATCAAGCGTAGTGGTTTTGGACGTGAACTTGGAGAATGG GGTATCCAGAATTACTTGAATATCAAGCAGGTGACTCAAGATATTTCTGATGAACCATGGGGATGGTACAAGTCTCCTTG AAAGCCGAATTCCAGCACACTGGCGGCCGTTACTAGATCCATCACACTGGCGGCCCGAACACGGAATTCAATGGAAGCAA TGATAAAAAAATACAAATAGAAAAGGAAAGGGAGGAAATACAAAAAAATAGAAGAGAAAAAGTCATACAAAGTTATATAC

## **FIG 36**

2. Sequence of gfp/BADH expression cassette (SEQ ID No. 2): CGGGCCCCCCCCGAGGTCGACGGTATCGATGAGCCTGATTATCCCTAAGCCCAATGTGAGTTTTCTAGTTGGATTTGC TCCCCCGCCGTCGTTCAATGAGAATGGATAAGAGGCTCGTGGGATTGACGTGAGGGGGCAGGGATGGCTATATTTCTGGG AGCGAACTCCGGGCGAATATGAAGCGCATGGATACAAGTTATGCCTTGGAATGAAAGACAATTCCGAATCCGCTTTGTCT ACCGGGAGACCACAACGGTTTCCCTCTAGAAATAATTTTGTTTAACTTTAAGAAGGAGATATACCCATGTCCATGAGTAA AGGAGAAGAACTTTTCACTGGAGTTGTCCCAATTCTTGTTGAATTAGATGGTGATGTTAATGGGCACAAATTTTCTGTCA TGGCCAACACTTGTCACTACTTTCTCTTATGGTGTTCAATGCTTTTCAAGATACCCAGATCATATGAAGCGGCACGACTT CTTCAAGAGCGCCATGCCTGAGGGATACGTGCAGGAGAGGACCATCTCTTTCAAGGACGACGAGGAACTACAAGACACGTG CTGAAGTCAAGTTTGAGGGAGACACCCTCGTCAACAGGATCGAGCTTAAGGGAATCGATTTCAAGGAGGACGGAAACATC TAACTTCAAAATTAGACACAACATTGAAGATGGAAGCGTTCAACTAGCAGACCATTATCAACAAAATACTCCAATTGGCG ATGGCCCTGTCCTTTTACCAGACAACCATTACCTGTCCACACAATCTGCCCTTTCGAAAGATCCCAACGAAAAGAGAGAC  ${\sf CACATGGTCCTTCTTGAGTTTGTAACAGCTGCTGGGATTACACATGGCATGGATGAACTATACAAATAATCTAGAAAGCC}$ CTTGGTTGACACGAGTATATAAGTCATGTTATACTGTTGAATAAAAAGCCTTCCATTTTCTATTTTGATTTGTAGAAAAC TAGTGTGCTTGGGAGTCCCTGATGATTAAATAAACCAAGATTTTCCATGGCGTTCCCAATTCCTGCTCGTCAGCTATTCA TCGACGGAGAGTGGAGAAACCCATTAAAAAAAAATCGCATACCCGTCATCAATCCGTCCACTGAAGAAATCATCGGTGAT ATTCCGGCAGCCACGGCTGAAGATGTGGAGGTTGCGGTGGCAGCTCGAAGAGCCTTTAGGAGGAACAATTGGTCAGC TGGAAACCATTGATCTGGGAAACCTTTTGATGAAGCAGTGCTGGACATTGATGACGTTGCTTCATGTTTTGAATATTTT GCCGGACAAGCAGAAGCTCTTGATGGTAAACAAAAGGCTCCAGTCACCCTGCCTATGGAAAGGTTCAAAAGTCATGTTCT CAGGCAGCCCTTGGTGTTGTTGGATTAATATCCCCATGGAATTACCCACTTCTAATGGCTACATGGAAAATTGCTCCAG CACTTGCTGCTGGGTGTACAGCTGTACTTAAGCCATCCGAGTTGGCATCTGTGACTTGTCTAGAATTCGGTGAAGTTTGC AACGAAGTGGGACTTCCTCCAGGCGTGTTGAATATCTTGACAGGATTAGGTCCAGATGCTGGTGCACCATTAGTATCACA CCCGATGTTGACAAGATTGCCTTTACTGGGAGTAGTGCCACTGGAAGCAAGGTTATGGCTTCTGCCCCAATTGGTTA AGCCTGTTACATTAGAACTTGGGGGTAAAAGTCCTATTGTAGTGTTTGAAGATGTTGATATTGATAAAGTTGTGGAATGG ACTATTTTTGGCTGTTTCTGGACAAATGGTCAAATATGTAGTGCAACGTCTAGACTGCTTGTGCATGAAAGTATTGCAGC  ${\sf TGAGTTTGTTGATAAGCTTGTAAAATGGACGAAAAACATTAAAATTTCTGACCCATTTGAAGAAGGATGCCGGCTTGGCC}$ CTGTTATTAGTAAAGGACAGTACGACAAAATTATGAAGTTCATATCAACAGCAAAGAGTGAGGGGGCAACTATTTTGTAT GGAGGTTCCCGTCCTGAGCATTTGAAGAAAGGTTATTACATTGAACCCACCATTGTAACTGATATCTCCACATCCATGCA AATATGGAAAGAGGAAGTTTTTGGCCCTGTCTTGTGTGTTAAAACATTTAGTTCCGAAGATGAAGCCATTGCATTGGCAA ATGATACAGAGTACGGTTTAGCTGCTGCTGTGTTTTCTAATGATCTTGAAAGATGTGAGAGGATAACGAAGGCTCTAGAA GTTGGAGCTGTTTGGGTTAATTGCTCACAACCATGCTTTGTTCAAGCTCCTTGGGGAGGCATCAAGCGTAGTGGTTTTTGG ACGTGAACTTGGAGAATGGGGTATCCAGAATTACTTGAATATCAAGCAGGTGACTCAAGATATTTCTGATGAACCATGGG GATGGTACAAGTCTCCTTGAAAGCCGAATTCCAGCACACTGGCGGCCGTTACTAGTGGATCCACTAGTAACGGCCGCCAG  ${\sf TTTTGTTTACATTATAGAAAAAGAAGGAGAGGTTATTTTCTTGCATTTATTCATGATTTGAGTATTCTATTTTGATTTTGT$ ATTTGTTTGGGCTGCGAGCT

## FIG 37

AATTTCGATGGATACAAGTTATGCCTTGGAATGAATTTCGGTTGATTAGGACTAGATCGTCGACGTAGAGAAGTCCGTAT CTGGGCACAACAGACAATCGGCTGCTCTGATGCCGCCGTGTTCCGGCTGTCAGCGCAGGGGCGCCCGGTTCTTTTGTCA GTCATCTCACCTTGCTCCTGCCGAGAAAGTATCCATCATGGCTGATGCAATGCGGCGGCTGCATACGCTTGATCCGGCTA GATCTGGACGAAGAGCATCAGGGGCTCGCGCCAGCCGAACTGTTCGCCAGGCTCAAGGCGCGCATGCCCGACGGCGATGA TCTCGTCGTGACCCATGGCGATGCCTGCTGCCGAATATCATGGTGGAAAAATGGCCGCTTTTCTGGATTCATCGACTGTG GCCGCTGGGTGTGGCGGACCGCTATCAGGACATAGCGTTGGCTACCCGTGATATTGCTGAAGAGCTTGGCGGCGAATGG GCTGACCGCTTCCTCGTGCTTTACGGTATCGCCGCTCCCGATTCGCAGCGCATCGCCTTCTATCGCCTTCTTGACGAGTT TTATAGAAAAAGAAGGAGGTTATTTCTTGCATTTATTCATGATTGAGTATTCTATTTTGATTTTGTATTTTGGG CTGCGAGCT



**FIG 38**